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(54) Title: MAINTENANCE OF MALE-STERILE PLANTS

(57) Abstract

Novel transgenic plants that have, stably integrated into their nuclear genome, a maintainer gene comprising a fertility-restorer gene and a pollen-lethality gene. The plants can be used to maintain a homogeneous population of male-sterile plants.

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MAINTENANCE OF MALE-STERILE PLANTS

This invention relates to a process for maintaining male-sterile plant lines that can be used for the production of hybrid seed of a crop, to maintainer plants that can be used in such a process, and to maintainer genes that can be used to produce such maintainer plants.

Background of the Invention

In many, if not most, plant species, the development of hybrid cultivars is highly desired because of their generally increased productivity due to heterosis: the superior performance of hybrid individuals compared with their parents (see, e.g., Fehr (1987) "Principles of Cultivar Development, Volume 1: Theory and Technique", MacMillan Publishing Company, New York; Allard (1960) "Principles of Plant Breeding", John Wiley and Sons, Inc., New York).

The development of hybrid cultivars of various plant species depends upon the capability to achieve almost complete cross-pollination between parents. This is most simply achieved by rendering one of the parent lines male-sterile (i.e., with pollen being absent or nonfunctional), for example, by manually removing the one parent's anthers or by providing the one parent with naturally occurring cytoplasmic or nuclear genes that prevent anther and/or pollen development and/or function, using classical breeding techniques (for a review of the genetics of malesterility in plants, see Kaul (1988) "Male Sterility in Higher Plants", Springer Verlag, New York).

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For hybrid plants where the seed is the harvested product (e.g., corn and oilseed rape), it is, in most cases, also necessary to ensure that fertility of the hybrid plants is fully restored. In plants in which the male-sterility is under genetic control, this requires the use of genes that can restore male-fertility. Hence, the development of hybrid cultivars is mainly dependent on the availability of suitable and effective sterility and restorer genes.

Endogenous nuclear loci are known for most plant species that contain genotypes which effect malesterility, and generally, such loci need to be homozygous for particular recessive alleles in order to result in a male-sterile phenotype. The presence of a dominant male-fertile allele at such loci results in male-fertility.

Recently, it has been shown that male-sterility can be induced in a plant by providing the plant with a nuclear male-sterility genotype that includes a chimaeric male-sterility gene comprising sequence (or male-sterility DNA) coding, for example, for a cytotoxic product (such as an RNase) and under the control of a promoter which is predominantly in selected tissue of the plant's male reproductive organs. In this regard, tapetum-specific promoters, such as the promoter of the TA29 gene of Nicotiana tabacum, have been shown to be particularly useful for this purpose (Mariani et al (1990) Nature 347:737; European patent publication 0,344,029). By providing the nuclear genome of the plant with such a male-sterility gene, an artificial nuclear male-sterility locus is created containing the artifical male-sterility genotype that results in a male-sterile plant.

In addition, it has been recently shown that male-fertility can be restored to such a nuclear male-sterile plant with a chimaeric fertilityrestorer gene comprising another DNA sequence (or fertility-restorer DNA) that codes, for example, for a protein that inhibits the activity of the cytotoxic product or otherwise prevents the cytotoxic product from being active at least in the selected tissue of the plant's male reproductive organs (EP 0,412,911). Bacillus gene of ` barnase the example, amyloliquefaciens codes for an RNase (Barnase) which can be inhibited by a protein (Barstar) that is encoded by the barstar gene of B. amyloliquefaciens. the barnase gene can be used construction of a chimaeric male-sterility gene while the barstar gene can be used for the construction of a chimaeric fertility-restorer gene. Experiments in different plant species (e.g., oilseed rape) have shown that such a chimaeric barstar gene can fully restore the male-fertility of male-sterile lines in which the male-sterility was due to the presence of such a chimaeric barnase gene (EP 0,412,911: Mariani al (1991) Proceedings of the CCIRC Rapeseed Congress, July 9-11, 1991 Saskatoon, Saskatchewan, Canada; Mariani et al (1992) Nature 357:384). coupling a marker gene, such as a dominant herbicide resistance gene (for example, the bar gene coding for (PAT) phosphinothricin acetyl transferase converts herbicidal phosphinothricin to a non-toxic compound [De Block et al (1987) EMBO J. 6:2513]), to male-sterility and/or fertility chimaeric the restorer gene, breeding systems can be implemented to select for uniform populations of male-sterile plants (EP 0,344,029; EP 0,412,911).

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The production of hybrid seed of any particular cultivar of a plant species requires the: maintenance of small quantities of pure seed of each inbred parent and 2) the preparation of quantities of seed of each inbred parent. Such larger quantities of seed would normally be obtained by several (usually two) seed-multiplication rounds, starting from a small quantity of pure seed ("basic seed") and leading, in each multiplication round, to a larger quantity of seed of the inbred parent and finally to a stock of seed of the inbred parent ("parent seed" or "foundation seed") which is of sufficient quantity to be planted to produce the desired quantities of hybrid seed. Of course, in each seed-multiplication round, larger planting areas (fields) are required.

In order to maintain and enlarge a small stock seeds of male-sterile plants, it necessary to cross the parent male-sterile plants with normal pollen-producing parent plants. offspring of such a cross will, in all cases, be a mixture of male-sterile and male-fertile plants, and the latter have to be removed from the former. male-sterile plants containing an artificial malesterility locus as described above, such removal can be facilitated by genetically linking the chimaeric male-sterility gene to a suitable marker gene, such as the bar gene, which allows the easy identification and removal of the male-fertile plants. EP 0,198,288 and US Patent 4,717,219, by comparison, describe methods for linking such marker genes (which can be visible markers or dominant conditional markers) to endogenous nuclear loci containing male-sterility genotypes.

However, even when suitable marker genes are linked to male-sterility genotypes, the maintenance of parent male-sterile plants still requires the removal from the field of a substantial number of For instance, in systems using a herbicide resistance gene (e.g., the bar gene) linked to a chimaeric male-sterility gene, only half of the parent stock will result in male-sterile plants, thus requiring the removal of the male-fertile plants by herbicide spraying prior to flowering. In any given field, the removal of male-fertile plants effectively reduces the potential yield of hybrid seed or the potential yield of male-sterile plants during each round of seed multiplication for producing of parent This is economically unattractive for many important crop species such as corn and oilseed rape. In order to minimize the number of male-fertile to be removed, male-fertile plants which have maintainer plants have been sought which, crossed with a male-sterile parent plant, produce a male-fertile offspring, minimum, preferably no, thereby minimizing or avoiding altogether the need to remove such male-fertile offspring. To solve an analogous problem, US Patents 3,710,511 and 3,861,079 described procedures for producing maintaining a homogenous population of male-sterile plants by using specific chromosomal abnormalities that are differentially transmitted to the egg and the sperm in the plants.

Summary of the Invention

In accordance with this invention, a cell of a transgenic plant ("the maintainer plant") is provided, in which the nuclear genome contains stably integrated therein: 1) at a first locus or malesterility locus, a male-sterility genotype in

homozygous condition; and 2) at a second locus or maintainer locus, a maintainer gene in heterozygous male-sterility locus the condition; maintainer locus preferably being unlinked; the maintainer gene being a foreign DNA sequence, a foreign sequence, chimaeric DNA preferably containing:

- a) a fertility-restorer gene that comprises at least:
 - i) a fertility-restorer DNA encoding a restorer RNA and/or protein or polypeptide which, when produced or overproduced in some or all of the cells, preferably stamen cells, of the plant, prevents phenotypic expression of the nuclear male-sterility genotype that would render the plant male-sterile in the absence of expression of the fertility-restorer DNA in the some or all stamen cells and
- ii) a restorer promoter capable of directing expression of the fertility-restorer DNA at least in the some or all of the cells, preferably stamen cells, of the plant, so that the phenotypic expression of the nuclear malesterility genotype is prevented, the fertility-restorer DNA being in the same transcriptional unit as, and under the control of, the restorer promoter and
- b) a pollen-lethality gene that is selectively expressed in microspores and/or pollen of the plant to produce nonfunctional pollen and that comprises at least:
- iii) a pollen-lethality DNA coding for a pollenlethality RNA and/or protein or polypeptide that, when produced or overproduced in the microspores and/or pollen, significantly

disrupts their metabolism, functioning and/or development and

iv) a pollen-specific promoter capable of directing expression of the pollen-lethality DNA selectively in the microspores and/or pollen of the plant, the pollen-lethality DNA being in the same transcriptional unit as, and under the control of, the pollen promoter.

The cell of the maintainer plant of this invention preferably also comprises, especially in the maintainer locus, at least one first marker gene which comprises at least:

- v) a first marker DNA encoding a first marker RNA and/or protein or polypeptide which, when present at least in a first specific tissue or specific cells of the plant, renders the plant easily separable from other plants which do not contain the first marker RNA, protein or polypeptide encoded by the first marker DNA at least in the first specific tissue or specific cells and
- vi) a first marker promoter capable of directing expression of the first marker DNA at least in the first specific tissue or specific cells, the first marker DNA being in the same transcriptional unit as, and under the control of, the first marker promoter.

The male-sterility genotype in the cell of the maintainer plant of this invention can be foreign or endogenous but is preferably a foreign, especially chimaeric, male-sterility gene which comprises:

1) a male-sterility DNA encoding a sterility RNA and/or protein or polypeptide which, when produced or overproduced in a stamen cell of the plant in the absence of the restorer RNA, protein or polypeptide,

significantly disturbs the metabolism, functioning and/or development of the stamen cell and

2) a sterility promoter capable of directing expression of the male-sterility DNA selectively in stamen cells of the plant, the male-sterility DNA being in the same transcriptional unit as, and under the control of, the sterility promoter.

The male-sterility genotype in the maintainer plant cell of this invention preferably comprises, especially in the male-sterility locus, at least one second marker gene which comprises at least:

- 3) a second marker DNA encoding a second marker RNA and/or protein or polypeptide which, when present at least in the second specific tissue or specific cells of the plant, renders the plant easily separable from other plants which do not contain the second marker RNA, protein or polypeptide encoded by the second marker DNA at least in the second specific tissue or specific cells and
- 4) a second marker promoter capable of directing expression of the second marker DNA at least in the second specific tissue or specific cells, the second marker DNA being in the same transcriptional unit as, and under the control of, the second marker promoter.

Also in accordance with this invention are provided the maintainer plants, the seeds of such plants, and plant cell cultures, all of which consist essentially of the cells of this invention.

Further in accordance with this invention are provided the maintainer gene and plasmids containing the maintainer gene, as well as bacterial host cells (e.g., <u>E. coli</u> or <u>Agrobacterium</u>) containing such plasmids.

Still further in accordance with this invention is provided a process for producing, preferably

enlarging, a homogeneous population of male-sterile plants and their seed that contain a nuclear male-sterility gene in homozygous condition, the process comprising the step of crossing the male-sterile plants with the maintainer plants of this invention. The seed from the resulting male-sterile plants can be harvested and grown into the male-sterile plants. Hybrid seed can then be produced by crossing the male-sterile plants with male-fertile plants of another inbred parent line used as pollinators.

Yet further in accordance with this invention is provided a process for producing, preferably enlarging, a population of the maintainer plants, comprising the step of selfing the maintainer plants.

Detailed Description of the Invention

A male-sterile plant of this invention is a plant of a given species with a nuclear male-sterility genotype.

A restorer plant of this invention is a plant of the same plant species containing, within its nuclear genome, a fertility-restorer gene that is able to restore the male-fertility in offspring which are obtained from a cross between the male-sterile plant and the restorer plant and which contain both the male-sterility genotype and the fertility-restorer gene.

A restored plant of this invention is a plant of the same species that is male-fertile and that contains, within its nuclear genome, the malesterility genotype and the fertility-restorer gene.

A parent plant or parent of this invention is a plant that can be used for the production of hybrid seed. The female or seed parent plant is the parent from which the hybrid seed is harvested. For the purposes of this invention, the female parent will

always be a male-sterile plant. The male or pollen parent is the parent that is used to fertilize the female parent. In many cases, the male parent will also be a restorer plant.

A line is the progeny of a given individual plant.

The male-sterility genotype of this invention is the genotype of at least one locus, preferably only one locus, in the nuclear genome of a plant (i.e., the male-sterility locus), the allelic composition of which can result in male-sterility in the plant. A male-sterility genotype can be endogenous to the plant, but it is generally preferred that it be foreign to the plant. Preferred foreign male-sterility genotypes are those in which the allele responsible for male-sterility contains a foreign male-sterility gene that comprises:

- 1) a male-sterility DNA encoding a sterility RNA and/or protein or polypeptide which, when produced or overproduced in a stamen cell of the plant, significantly disturbs the metabolism, functioning and/or development of the stamen cell and
- 2) a sterility promoter capable of directing expression of the male-sterility DNA selectively in stamen cells of the plant, the male-sterility DNA being in the same transcriptional unit as, and under the control of, the sterility promoter.

Such a male-sterility gene is always a dominant allele at a foreign male-sterility locus. The recessive allele corresponds to the absence of the male-sterility gene in the nuclear genome of the plant.

male-sterility foreign Preferred sterility promoters that can be used in the malefemale parent plants sterility genes in this invention maintainer plants of described in EP 0,344,029. A particularly useful male-sterility DNA codes for Barnase (Hartley (1988) J.Mol. Biol. 202:913). Particularly useful sterility promoters are tapetum-specific promoters such as: the promoter of the TA29 gene of Nicotiana tabacum (EP 0,344,029) which can be used in tobacco, oilseed rape, lettuce, chicory, corn and other plant species; the PT72, the PT42 and PE1 promoters from rice, the sequences of which are given in SEQ ID no. 7, SEQ ID no. 8, and SEQ ID no. 9, respectively, of the Sequence Listing and which can be used in rice and species (PCT application plant PCT/EP 92/00274); and the PCA55 promoter from corn, the sequence of which is given in SEQ ID No. 10, which can be used in corn and other plant species (PCT application PCT/EP 92/00275).

A preferred endogenous male-sterility genotype in which a recessive allele ("m") is one homozygous condition (m/m) at a male-sterility locus produces male-sterility. At a male-sterility locus, male-fertility would otherwise be encoded by a corresponding dominant allele ("M"). Such a malesterility genotype is known in many plant species (see Kaul (1988) supra; and 1992 issues of Maize Genetics Cooperation Newsletter, published by the Department of Agronomy U.S. Department and University Of Missouri, Columbia, Agriculture, Missouri, U.S.A.). The DNA sequences in the nuclear genome of a plant corresponding to m and M alleles identified by gene tagging, insertional mutagenesis using transposons,

means of T-DNA integration (see, e.g., Wienand and Saedler (1987) In "Plant DNA Infectious Agents", Ed. by T.H.Hohn and J.Schell, Springer Verlag, New York, p. 205; Shepherd (1988) In "Plant Molecular Biology: a Practical Approach", IRL Press, p. 187; Teeri et al (1986) EMBO J. 5:1755).

Fertility-restorer DNAs and restorer promoters that can be used in the maintainer genes of this invention with a foreign male-sterility genotype have been described in EP 0,412,911. In this regard, fertility-restorer genes in which the fertility-restorer DNA encodes Barstar (Hartley (1988) J.Mol. Biol. 202:913) and is under control of tapetum-specific promoters, such as those described above as sterility promoters, are of particular use. In particular, it is believed that a fertility-restorer DNA coding for a mutant of Barstar, in which the cysteine residue at its position 40 is replaced by serine (Hartley (1989) TIBS 14:450), functions better in restoring the fertility in the restored plants of some species.

When an endogenous male-sterility genotype is homozygous for a recessive allele m, it is preferred that the fertility-restorer gene be the dominant allele M of that male-sterility genotype, preferably under the control of its own promoter. The DNA corresponding to such a dominant allele, including its natural promoter, can be isolated from the nuclear genome of the plant by means of gene tagging as described above.

The pollen-lethality DNAs that are used in the pollen-lethality genes of this invention preferably encode an RNA and/or a protein or polypeptide that, when expressed in microspores or pollen, significantly disrupts their metabolism, functioning

In this regard, the pollenand/or development. encode RNAs, proteins DNAs can polypeptides such as are encoded by the malein EP 0,344,029. described DNAs sterility particular interest are male-sterility DNAs that encode ribonucleases (EP 0,344,029) such as RNase T1 from Aspergillus oryzae (Quaas et al (1988) Eur. J. Barnase from Bacillus 173:617) or Biochem. (1988)J.Mol.Biol. amyloliquefaciens (Hartley 202:913).

So that the pollen-lethality DNA is expressed pollen selectively in microspores or maintainer plant, it is preferred that the pollenspecific promoter, which the controls lethality DNA in the pollen-lethality gene, be a gene expression capable of directing promoter selectively in the microspores and/or pollen of the plant. Such a pollen-specific promoter can be an endogeneous promoter or a foreign promoter and can be from the nuclear genome or from the mitochondrial or chloroplast genome of a plant cell, but in any event, the pollen-specific promoter is foreign in nuclear genome of the plant being transformed. Preferably the pollen-specific promoter causes the pollen-lethality DNA to be expressed only in the microspores and/or pollen, i.e., after meiosis of the microsporocytes in the anthers. The pollen-specific promoter can be selected and isolated in a well known manner from a plant species, preferably the plant species to be rendered male-sterile, so that the pollen-specific promoter directs expression of the pollen-lethality DNA selectively in the microspores as to kill or disable and/or pollen SO in which the microspores and/or pollen lethality gene is expressed. The pollen-specific promoter is preferably also selected and isolated so that it is effective to prevent expression of the pollen-lethality DNA in other tissues of the plant. For example, a suitable endogeneous pollen-specific promoter can be identified and isolated in a plant, to be rendered male-sterile, by:

- searching for an mRNA which is only present in the plant during the development of its microspores and/or pollen;
- optionally isolating the microspore- and/or pollen-specific mRNA;
- 3. preparing and isolating a cDNA from the microspore- and/or pollen-specific mRNA;
- 4. using this cDNA as a probe to identify regions in the plant genome which contain DNA coding for the corresponding microspore- and/or pollenspecific DNA or alternatively using inverse polymerase chain reactions for the geometric amplification of the DNA sequences which flank, upstream and downstream, a chosen core region of the genomic DNA corresponding to the sequence of the microspore- and/or pollen-specific cDNA; and
- 5. identifying the portion of the plant genome that is upstream (i.e. 5') from the DNA coding for the microspore- and/or pollen-specific mRNA and that contains the promoter of this DNA.

Examples of such pollen-specific promoters are well known (see MacCormick (1991) TIG 7:298). In this regard, Hamilton et al (1989) Sex. Plant Reprod. 2:208 describes a pollen-specific clone ("Zmg13") from maize inbred line W-22, and the use of the promoter sequences of the clone to direct pollen-specific expression in tobacco has been described by Guerrero et al (1990) Mol.Gen.Genet. 224:161). Other pollen-specific promoters that are likewise believed

to be useful are: the promoter of the gene corresponding to the <u>Nicotiana tabacum</u> pollen-specific cDNA NTPc303 described by Weterings et al (1992) Plant Mol. Biol. <u>18</u>:1101; and the promoter of the gene corresponding to the <u>Brassica napus</u> pollen-specific cDNA B54 described by Shen and Hsu (1992) Mol. Gen. Genet. 234:379.

If the fertility-restorer DNA in the fertilityrestorer gene of the maintainer gene is expressed in microspores and/or pollen at the same time as the pollen-lethality DNA is expressed (due for instance to the activity of the restorer promoter in microspores and/or pollen), it is preferred that the pollen-lethality DNA be different from the malesterility DNA (the expression of which is intended to be prevented by expression of the fertility-restorer DNA of the maintainer gene). For example, if the male-sterility DNA encodes Barnase in the malesterile plants to be maintained, the fertilityrestorer DNA in the maintainer gene should encode Thus, if the restorer promoter in Barstar. gene also directs expression of maintainer fertility-restorer DNA in microspores and/or pollen and at the same time as the pollen-lethality DNA is pollen-lethality DNA preferably expressed, the should not encode Barnase but rather, for example, another RNAse such as RNAse T1.

First and second marker DNAs and first and second marker promoters that can be used in the first and second marker genes of this invention are also well known (EP 0,344,029; EP 0,412,911). In this regard, it is preferred that the first and second marker DNAs be different, although the first and second marker promoters may be the same.

The fertility-restorer gene, the male-sterility gene, the pollen-lethality gene, and the first and second marker genes in accordance with this invention generally foreign DNA sequences, preferably foreign chimaeric DNA sequences. Such foreign DNA sequences are preferably provided with suitable 3' transcription regulation sequences polyadenylation signals, downstream (i.e. 3') from respective fertility-restorer DNA, malesterility DNA, pollen-lethality DNA, and first and second marker DNAs. In this regard, either foreign or endogenous, transcription termination polyadenylation signals suitable for obtaining expression of such DNA sequences can be used. For example, the foreign 3' untranslated ends of genes, such as gene 7 (Velten and Schell (1985) Nucl. Acids Res. 13:6998), the octopine synthase gene (De Greve et al (1982) J.Mol. Appl. Genet. 1:499; Gielen et al (1983) EMBO J. 3:835; Ingelbrecht et al (1989) The Plant Cell 1:671) the nopaline synthase gene of the T-DNA region of Agrobacterium tumefaciens Tiplasmid (De Picker et al (1982) J.Mol. Appl. Genet. 1:561), the chalcone synthase gene (Sommer and Saedler (1986) Mol. Gen. Genet. 202: 429-434), and the CaMV 195/355 transcription unit (Mogen et al (1990) The Plant Cell 2:1251-1272), can be used.

By "foreign" with regard to a gene or genotype of this invention is meant that the gene or genotype contains a foreign DNA sequence such as a malesterility DNA, a fertility-restorer DNA, a pollen-lethality DNA, or a marker DNA and/or a foreign promoter such as a sterility promoter, a restorer promoter, a pollen-specific promoter or a marker promoter. By "foreign" with regard to any DNA sequence, such as a coding sequence or a promoter, in a gene or genotype of this invention is meant that such a DNA is not in the same genomic environment in a plant cell, transformed with such a DNA in

accordance with this invention, as is such a DNA when it is naturally found in the cell of the plant, bacteria, animal, fungus, virus or the like, from which such a DNA originates. This means, for example, that a foreign fertility-restorer DNA, male-sterility DNA, pollen-lethality DNA, or marker DNA can be: 1) a nuclear DNA in a plant of origin; 2) endogenous to the transformed plant cell (i.e., from a plant of origin with the same genotype as the plant being transformed); and 3) within the same transcriptional unit as its own endogenous promoter and 3' end transcription regulation signals (from the plant of origin) in the foreign gene or genotype in the transformed plant cell; but 4) inserted in the nuclear different place genome transformed plant cell than it was in the plant of is not so that it surrounded origin transformed plant cell by the genes which surrounded it naturally in the plant of origin. Likewise, a foreign fertility-restorer DNA, male-sterility DNA, pollen-lethality DNA, or marker DNA can also, for example, be: 1) a nuclear DNA in a plant of origin; and 2) endogenous to the transformed plant cell; but 3) in the same transcriptional unit as a different (i.e., not its own) endogenous promoter and/or 3' end transcription regulation signals in chimaeric gene or genotype of this invention in a transformed plant cell. A foreign fertility-restorer DNA, male-sterility DNA, pollen-lethality DNA, or marker DNA can also, for example, be: 1) a nuclear DNA in a plant of origin; and 2) endogenous to the transformed plant cell; but 3) in the transcriptional unit as a heterologous and/or 3' end transcription regulation signals in a foreign chimaeric gene or genotype of this invention in a transformed plant cell. A foreign fertilityrestorer DNA, a male-sterility DNA, pollen-lethality
DNA, or marker DNA can also, for example, be
heterologous to the transformed plant cell and in the
same transcriptional unit as an endogenous promoter
and/or 3' transcription regulation signals (e.g.,
from the nuclear genome of a plant with the same
genotype as the plant being transformed) in a foreign
chimaeric DNA sequence of this invention in a
transformed plant cell. Preferably, each fertilityrestorer DNA, male-sterility DNA, pollen-lethality
DNA, and marker DNA of this invention is heterologous
to the plant cell being transformed.

By "heterologous" with regard to a DNA, such as a fertility-restorer DNA, a male-sterility DNA, a pollen-lethality DNA, a marker DNA, a fertilityrestorer promoter, a sterility promoter, a pollenspecific promoter or a marker promoter or any other DNA sequence in a gene or a genotype of this invention is meant that such a DNA is not naturally found in the nuclear genome of cells of a plant with the same genotype as the plant being transformed. Examples of heterologous DNAs include chloroplast and mitochondrial DNAs obtained from a plant with the same genotype as the plant being transformed, but preferred examples are chloroplast, mitochondrial, and nuclear DNAs from plants having a different genotype than the plant being transformed, DNAs from animal and bacterial genomes, and chromosomal and plasmidial DNAs from fungal, bacterial and viral genomes.

By "chimaeric" with regard to a foreign DNA sequence of this invention is meant that at least one of its coding sequences: 1) is not naturally found under the control of the promoter present in the

foreign DNA sequence; and/or 2) is not naturally found in the same genetic locus as at least one of its associated marker DNAs. Examples of foreign chimaeric DNA sequences of this invention comprise: a pollen-lethality DNA of bacterial origin under the control of a pollen-specific promoter of plant origin; and a pollen-lethality DNA of plant origin under the control of a pollen-specific promoter of plant origin and in the same genetic locus as a marker DNA of bacterial origin.

By "endogenous" with respect to a gene or genotype of this invention is meant that it is not foreign.

genotypes of this foreign genes and The the male-sterility gene and invention. such as fertility-restorer gene the and genotype, the pollen-lethality gene, can be described like any other genotype: capital letters denote the presence of the foreign genes and genotypes (the dominant allele) while small letters denote their absence (the recessive allele). Hence, in this description of the invention, "S" and "s" will denote the respective presence and absence of the male-sterility gene, "R" and "r" will denote the respective presence absence of the fertility-restorer gene, and "P" and "p" will denote the respective presence and absence of the maintainer gene.

For an endogeneous male-sterility genotype of this invention, "m" will denote the recessive allele, and "M" will denote the dominant allele. Thus, the recessive allele m in homozygous condition (m/m) at a male-sterility locus would result in male-sterility, and the dominant allele M, when present at a male-sterility locus either in homozygous or heterozygous condition, results in male-fertility.

The cell of a plant, particularly a plant capable of being infected with Agrobacterium such as most dicotyledonous plants (e.g. Brassica napus), can be transformed using a vector that is a disarmed Tiplasmid containing the male-sterility gene and/or the fertility-restorer gene and/or the pollen-lethality gene and/or the maintainer gene and/or the marker invention and carried gene(s) this of Agrobacterium. This transformation can be carried out using the procedures described, for example, in EP 0,116,718 and EP 0,270,822. Preferred Ti-plasmid vectors contain a foreign DNA sequence of this invention between the border sequences, or at least located to the left of the right border sequence, of the T-DNA of the Ti-plasmid. Of course, other types of vectors can be used to transform the plant cell, using procedures such as direct gene transfer (as described, for example, in EP 0, 233,247), pollen mediated transformation (as described, for example, in EP 0,270,356, PCT publication WO 85/01856, and US virus-mediated patent 4,684,611), plant RNA transformation (as described, for example, EP 0,067,553 and US patent 4,407,956) and liposomemediated transformation (as described, for example, in US patent 4,536,475). Cells of monocotyledonous plants, such as the major cereals including corn, rice, wheat, barley and rye, can be transformed as described in PCT application PCT/EP 91/02198. In case the plant to be transformed is corn, other recently developed methods can also be used such as, for example, the methods described for certain lines of corn by Fromm et al (1990) Bio/Technology 8:833, Gordon-Kamm et al (1990) Bio/Technology 2:603 and Gould et al (1991) Plant Physiol. 95:426. In case the plant to be transformed is rice, recently developed

methods can also be used such as, for example, the method described for certain lines of rice by Shimamoto et al (1989) Nature 338:274, Datta et al (1990) Bio/Technology 8:736 and Hayashimoto et al (1990) Plant Physiol. 93:857.

The so-transformed cell can be regenerated into a mature plant, and the resulting transformed plant can be used in a conventional breeding scheme to produce more transformed plants with characteristics or to introduce the male-sterility fertility-restorer gene, the pollenand/or marker genes lethality gene, the maintainer gene of this invention in other varieties of the same or related plant species. Seeds obtained such plants contain the gene(s) of this invention as a stable genomic insert.

The maintainer plant of this invention is of the same species as a male-sterile plant line and can be used for the maintenance of the male-sterile line, i.e. to maintain a homogeneous population of male-sterile plants and a stock of pure seed of the female parent. The maintainer plant of this invention is itself a plant in which male-fertility has been restored and the genome of which contains both a male-sterility genotype and, in the maintainer locus, a fertility-restorer gene of this invention.

If a plant line with a homozygous male-sterility genotype $(A^{m/m}$ or $A^{S/S})$ is available, a maintainer plant for the male-sterile line can be directly obtained by transforming a male-sterile plant of the line with the maintainer gene of this invention and then selecting those transgenic plants which are male-fertility restored plants and in which the maintainer gene is stably integrated in the nuclear genome so that the genetic locus of the male-

sterility genotype and of the maintainer gene are unlinked and segregate independently.

If the male-sterility genotype is foreign to the plant line, alternative strategies can be followed. For example, the maintainer plant of the present invention can be obtained by: transforming cell of the plant line (A) with the maintainer gene of this invention (P); and then regenerating, from the so-transformed plant cell, a transgenic plant containing, stably integrated in its genome, the (A^{P/P}) can maintainer gene. Such a transgenic plant then be crossed as a female parent with a plant A^{S/s,R/r} of the same line, which contains at separate loci in its genome a male-sterility gene (S) and a corresponding fertility-restorer gene (R), both in lacks heterozygous condition, but which maintainer gene. Thus, the cross is in fact: AS/s,R/r,p/p (male) $x A^{s/s,r/r,p/p}$ (female), and the offspring with the genotype A^{S/s,r/r,P/p} (or hereinafter "A^{S/s,P/p}" for convenience) are selected and selfed. One eighth of the offspring that have the desired genotype $(A^{S/S,P/P})$ for a maintainer plant of this invention can then be selected. Another eighth of the offspring with the genotype (AS/S,p/p) can be used as male-sterile plants to be maintained.

Isolation of plants with desired genotypes can be achieved by means of conventional testcrosses Fehr (1987)supra), preferably e.q., supplemented by detection of the presence of specific at the DNA level, e.g., by means amplification of DNA fragments by the polymerase chain reaction, by Southern blot analysis and/or by phenotypic analysis for the presence and expression of first or second marker genes of this invention.

The cross of a male-sterile plant containing a male-sterility genotype in homozygous condition ($A^{S/S}$ or $A^{m/m}$) with a maintainer plant of this invention ($A^{S/S,P/P}$ or $A^{m/m,P/P}$, respectively) results in a population of seeds that all contain the male-sterility genotype in homozygous condition ($A^{S/S}$ or $A^{m/m}$, respectively) because the maintainer gene is not transmitted through the pollen. This property can be used to advantage in maintaining the basic seed and in the multiplication of basic seed for the final production of parent seed.

The maintainer plants of this invention (ASS,P/P or A^{m/m,P/p}) can themselves be maintained by selfing. The offspring of such selfing will consist of 50% male-fertile maintainer plants $(A^{S/S,P/p} \text{ or } A^{m/m,P/p},$ respectively) and 50% male-sterile plants containing the male-sterility genotype in homozygous condition $(A^{S/S} \text{ or } A^{m/m}, \text{ respectively })$. If desired, the malesterile plants can be removed either manually on the basis of the male-sterile phenotype or, if maintainer gene comprises a suitable first marker gene, preferably a first marker gene whose expression confers herbicide resistance to the plant, by using the phenotypic expression of the first marker gene (e.g, by applying herbicide to the offspring so that male-sterile plants that lack the resistance gene are killed while maintainer plants with the herbicide-resistance gene survive).

Thus, the maintainer plant of this invention can be easily used to maintain a homogeneous population of male-sterile plants. In this regard, basic seed of a female parent of a given plant species can be crossed with an appropriate male parent to produce hybrid seed. Also, the maintainer plant of this invention can be used economically to multiply the

basic seed of a female parent of a given plant species, so as to obtain sufficient quantities of female parent seed that can be crossed with an appropriate male parent to produce desired quantities of hybrid seed.

A male-sterile line, that is maintained and multiplied by the use of the maintainer plants of this invention, can be used for the production of hybrid seed. In principle, the male-sterile line (A^{S/S}) can be crossed directly with another male parent line (B^{s/s}) to produce hybrid seed (AB^{\$/s}). However, as all hybrid plants are male-sterile, no reproduction and no seed set will occur. This is not a problem if the seed is not the harvested product (e.g., with lettuce), but where seed is the harvested product (e.g., with corn and oilseed rape), male-fertility in the hybrid plants should be at least partially restored. This can be accomplished by crossing the male-sterile line with a male-fertile parent line (e.g., BR/R) that is also a restorer line, i.e. that also contains a fertility-restorer gene (R). The hybrids produced $(AB^{S/s,R/r})$ will be fully male-fertile. Alternatively the male-sterile-line (AS/S) can first be crossed with the male-fertile line (As/s) just prior to hybrid seed productions. This has the advantage of giving a further multiplication of the female parent line. The offspring (AS/s) can then be crossed with a suitable male-fertile parent line (Bs/s,r/r) to produce hybrid seed that is 50% malefertile. If hybrid seed with 100% male fertility is desired, the offspring can be crossed with a suitable restorer male parent line (Bs/s,R/R).

In the case of a male-sterile line in which male-sterility is due to an endogeneous male-sterility genotype $(A^{m/m})$ at a male-sterility locus,

hybrid seed can easily be produced by crossing the male-sterile line $(A^{m/m})$ with a line that is homozygous with respect to the endogenous dominant (male-fertility) allele at that male-sterility locus $(B^{M/M})$. All hybrid offspring of this cross will have the genotype $AB^{M/m}$ and will be fertile.

The maintainer plants of this invention can also be used as pollinator (i.e., male-fertile) plants in a cross with wild-type plants $(A^{s/s,p/p})$ of the same inbred line. The progeny of this cross will all be male-sterile and heterozygous for the male-sterility genotype $(A^{s/s,p/p})$. The progeny can therefore be used directly for hybrid seed production by crossing with a pollinator plant line B $(B^{s/s,p/p})$. This scheme only requires a male-sterilization of the wild-type plants, for example by manually removing the anthers (e.g., in corn) or by using a male gametocide.

Of course, by using the maintainer plants of this invention to maintain a homogeneous population plants that are homozygous with respect to a male-sterility allele (whether dominant or recessive) that is encoded in the nuclear genome, the maintainer plants acquire many of the characteristics of plants of a cytoplasmic male-sterile line. However, plants do not have one of the major disadvantages of cytoplasmic male-sterile plants, namely cytoplasmic uniformity of the various male-sterile lines which, in corn, has led to serious problems (see Craig (1977) In "Corn and Corn Improvement", G.F. Sprague, ed., American Society of Agronomy, Inc., Publisher, p. 671).

Thus, the maintainer gene of this invention, when introduced into a particular line of a plant species, can always be introduced into any other line by backcrossing, but since the maintainer gene can

only be transmitted through an egg, it will always be associated with the cytoplasm of the line in which it was initially introduced. However, since a maintainer plant line is only used for maintenance of a malesterile line and not as a female parent for hybrid seed production, the hybrid seed will always contain the cytoplasm of the female parent, as desired.

illustrate this following Examples The indicated. Unless otherwise invention. experimental procedures for manipulating recombinant DNA were carried out by the standardized procedures "Molecular in Sambrook et al (1989)Cloning: A Laboratory Manual, Second Edition", Cold Spring Harbor Laboratory Press, N.Y. polymerase chain reactions ("PCR") were performed using the conditions, under conventional polymerase (Cat. No. 254L - Biolabs New England, 01915, U.S.A.) isolated MA Beverley, Thermococcus litoralis (Neuner et al (1990) Arch. Oligonucleotides Microbiol. 153:205-207). by the methods described by Kramer and designed in Enzymology 154:350 Methods Fritz (1968) synthesized by the phosphoramidite method (Beaucage and Caruthers (1981) Tetrahedron Letters 22:1859) on an applied Biosystems 380A DNA synthesizer (Applied Biosystems B.V., Maarssen, Netherlands).

The following bacterial strains and plasmids, used in the Examples, are available from the Deutsche Sammlung für Mikroorganismen und Zellkulturen ("DSM"), Mascheroder Weg 1B, Braunschweig, Germany:

Bacterial strain	plasmid	DSM No	Date of
			Deposit
E. coli WK6	pMa5-8	DSM 4567	May 3, 1988
E. coli WK6	pMc5-8	DSM 4566	May 3, 1988

In the Examples, reference will be made to the following Figure and Sequence Listing:

Figure

Figure 1: Ten-step procedure to obtain corn (e.g. H99) maintainer plants of the invention

Sequence Listing

genomic DNA comprising the promoter of SEQ ID no. 1: the Zml3 gene from Zea mays sequence of plasmid "pVE144" SEQ ID no. 2: SEQ ID no. 3: sequence of plasmid "pVE108" SEQ ID no. 4: sequence of oligonucleotide "MDB80" SEQ ID no. 5: sequence of oligonucleotide "MDB81" SEQ ID no. 6: sequence of oligonucleotide "MDB82" comprising the SEQ ID No. 7: genomic DNA specific promoter "PT72" from rice comprising the genomic DNA SEQ ID No. 8: specific promoter "PT42" from rice comprising the anther genomic DNA SEQ ID No. 9: specific promoter "PE1" from rice DNA comprising the anther SEQ ID No. 10: genomic specific promoter "PCA55" from corn SEQ ID No. 11: Oligonucleotide Zm13OLI2 SEQ ID No. 12: Oligonucleotide Zm13OLI1 SEQ ID No. 13: Oligonucleotide Zml3OLI5 SEQ ID No. 14: Oligonucleotide BXOL2 SEQ ID No. 15: Oligonucleotide TA29SBXOL2 SEQ ID No. 16: Oligonucleotide PTA290L5 SEQ ID No. 17: EcoRI-HindIII fragment of pTS218 carrying

the maintainer gene.

Examples

Example 1: Isolation of the pollen-specific promoter of the Zml3 gene from maize.

A pollen-specific cDNA from Zea mays inbred line W-22, designated as "Zmc13", has been isolated and characterized by Hanson et al (1989) The Plant Cell 1:173. The corresponding genomic clone, designated as "Zmg13", containing substantial portions of the 5' flanking region has been isolated and characterized by Hamilton et al (1989) Sex. Plant Reprod. 2:208 (see also Hamilton et al (1992) Plant Mol. Biol. 18:211). The complete sequence of Zmg13 is shown in SEQ ID no. 1, and its promoter region will hereinafter be referred to as the "Zm13 promoter".

A corresponding promoter region from corn inbred line H99 was isolated as follows. Genomic DNA of was prepared as described inbred line H99 Dellaporta et al (1983) Plant Mol. Biol. 1:19-21. Using the genome as a substrate, a 1471 bp PCR using was amplified by the fragment oligonucleotides MDB80 and MDB82, the sequences of which are shown in SEQ ID no. 4 and SEQ ID no. 6, respectively. MDB80 corresponds to nucleotides 8 to Zmg13, while MDB82 is complementary nucleotides 1458 to 1478 of Zmg13. Then, the purified amplified 1471 bp fragment was used as a substrate for the amplification by PCR of a 1422 bp fragment, using the oligonucleotides MDB80 and MDB81. MDB81 is complementary to nucleotides 1409 to 1429 of Zmg13, and its sequence is shown in SEQ ID no. 5. using MDB81, a NcoI site is created in the amplified 1422 bp fragment at the ATG translation initiation

The 1422 bp fragment is then ligated in an Smal site of pGEM2 (Promega Corporation, Madison,

Wisconsin 53711, U.S.A.), yielding plasmid pMDB13, and the fragment is sequenced (Maxam and Gilbert (1980) Meth. Enzymol. 65:499). The pollen-specific promoter of the Zm13 gene of corn inbred line H99 is obtained from pMDB13 as a EcoRV-NcoI fragment.

The Zm13 promoter is also cloned as follows. Genomic DNA of Zea mays line H99 is prepared as Using the genomic DNA described above. substrate, the following two fragments are amplified by means of PCR: 1) a 875 bp fragment is amplified using the oligonucleotides MDB80 (SEQ ID No. 4) and ZM130LI2 (which is complementary to nucleotides 859 to 882 of Zmg13 and which sequence is given in SEQ ID No. 11); and 2) a 661 bp fragment is amplified using the oligonucleotides Zml3OLI1 (which corresponds to nucleotides 767 to 791 of Zmg13 and which sequence is given in SEQ ID No. 12) and Zm13OLI5 (which is partially complementary to nucleotides 1397 to 1423 of Zmg13 and which sequence is given in SEQ ID No. The 875 bp fragment, corresponding to the upstream region of the Zm13 promoter, is cloned into the SmaI site of pGEM2, yielding plasmid pTS204. 661 bp fragment, corresponding to the downstream region of the Zml3 promoter, is digested with NcoI and cloned into plasmid pJB66 (Botterman and Zabeau (1987) DNA 6:583) digested with EcoRV and NcoI, yielding plasmid pTS203. Both fragments partly overlap and share a BstXI site in the region of overlap. Ligation of the 567 bp EcoRV-BstXI fragment of pTS204 and the 638 bp BstXI-NcoI fragment of pTS203 results in a 1205 bp fragment corresponding to the Zm13 promoter. This 1205 bp fragment, as cloned from line H99, is sequenced, and its sequence is found to be identical to the corresponding fragment of Zmgl3 from line W-22 as given in SEQ ID No.1 except at position 276 (G in W-22 is T in H99), 410 (G in)

W-22 is A in H99), and 1205-1206 (GC in W-22 is GGC in H99, thus corresponding to a 1 nucleotide insertion), numberings being as in SEQ ID No. 1.

Example 2: Construction of plant transformation vectors comprising a maintainer gene that contain DNA encoding Barstar under the control of the TA29 promoter and DNA encoding Barnase under the control of the Zm13 promoter.

The 1205 bp EcoRV-NcoI fragment of pMDB13 is ligated to the large EcoRI-SmaI fragment of plasmid pVE144 and to the 739 bp EcoRI-NcoI fragment of pVE108, yielding plasmid pGSJVR1. Plasmid pVE144, the sequence of which is shown in SEQ ID no. 2, is a plasmid derived from plasmid pUC18 (Yanisch-Perron et al (1985) Gene 33:103) and containing DNA encoding neomycin phosphotranferase (neo) under the control of the 35S3 promoter (EP 0,359,617) from Cauliflower Mosaic Virus isolate CabbB-JI (Hull and Howell (1978) Virology <u>86</u>:482) and DNA encoding the (Hartley (1988) J.Mol.Biol. 202:913) under control of the tapetum-specific promoter of the TA29 gene of Nicotiana tabacum (EP 0,344,029; Seurinck et (1990) Nucleic Acids Res. <u>18</u>:3403). pVE108, the sequence of which is shown in SEQ ID no. 3, is a plasmid derived from pUC18 and containing DNA encoding phosphinothricin acetyl transferase (bar) (EP 0,242,236) under the control of the 35S3 promoter and DNA encoding Barnase (Hartley (1980) supra) under the control of the TA29 promoter. The resulting plasmid, pGSJVR1 (which is subsequently renamed "pTS210"), is a pUC18-derived plasmid that contains a maintainer gene of this invention comprising: DNA encoding Barnase as the pollen-lethality DNA, the Zml3 promoter as the pollen-specific promoter, DNA

encoding Barstar as the fertility-restorer DNA, the TA29 promoter as the restorer promoter, neo as the first marker DNA and the 35S3 promoter as the first marker promoter.

pTS210 is also obtained as follows. The 0.9 kb BstXI-SacI fragment of pTS204 is ligated to the large BstXI-SacI fragment of pTS203, yielding plasmid pTS206. The 1.47 BglII-NcoI kb fragment ofpTS206 is then ligated to the large NcoI-BglII fragment of pVE108, yielding plasmid pTS207. Finally, the 1.9 kb EcoRV-Eco-RI fragment of pTS207 is ligated to the large Eco-RI-SmaI fragment of pVE144, yielding plasmid pTS210.

A plasmid pTS218, which differs from pTS210 by carrying the <u>bar</u> gene as a selectable marker gene, is also obtained as follows:

- a 255 bp DNA fragment, designated as bxx and carrying the translation initiation site of the PTA29-barstar gene, is obtained by PCR using pVE144 as a template and oligonucleotides BXOL2 (SEQ ID No. 14) and TA29SBXOL2 (SEQ ID No. 15) as primers.
- a 492 bp DNA fragment is prepared by PCR using pVE108 and bxx as a template and oligonucleotides PTA290L5 (SEQ ID No. 16) and BXOL2 as primers. This 492 bp fragment is digested with AsnI and BspEI, and a 274 bp fragment is purified on gel and ligated to the 6.28 kb fragment of pVE144 which was digested with BspEI and partially digested with AsnI. The resulting plasmid is designated as pVEK144 and carries the PTA29-barstar-3'nos chimeric gene

with an optimized translational initiation context.

- pVEK144 is digested with MunI and HindIII, and the 3.7 kbp fragment is isolated and ligated to the 1.7 kbp MunI-HindIII fragment of pVE108, yielding plasmid pVEB144 which carries the PTA29-barstar-3'nos and the P35S-bar-3'nos chimeric genes.
- the EcoRI-HindIII fragment of pVEB144, containing the two chimeric genes, is ligated to the large EcoRI-HindIII fragment of pUCNew2, yielding plasmid pVEC144. pUCNew2 is derived from pUC19 as described in WO 92/13956.
- finally, the large EcoRI-SmaI fragment of pVEC144 is ligated to the 1.9 bp EcoRV-EcoRI fragment of pTS207, yielding plasmid pTS218.

Plasmid pTS218 carries three chimeric genes, i.e., PTA29-barstar-3'nos (with optimized translational initiation context), P35S-bar-3'nos, and PZM13-barnase-3'nos. The EcoRI-HindIII fragment of pTS218 carrying these three chimeric genes is presented in the sequence listing as SEQ ID No. 17.

steps of vector construction involving fragments of the barnase DNA, such as pVE108, pVE144, and ptS210, are carried out in E. coli strain MC1061 containing the cointegrate plasmid R702::pMc5BS which is obtained as follows. Plasmid pMc5BS, containing the barstar gene (encoding an inhibitor of barnase) under the control of the tac promoter (De Boer et al Proc. Acad. Sci. USA 80:21), (1983)Natl. constructed by: cloning the EcoRI-HindIII fragment of plasmid pMT416 (Hartley (1988) supra) into the EcoRI and HindIII sites of plasmid pMc5-8 (DSM 4566); and starting deleting the sequence initiation codon of the phoA signal sequence and ending with the last nucleotide before the

translation initiation codon of the barstar-coding region by means of a looping-out mutagenesis procedure as generally described by Sollazo et al (1985) Gene 37:199.

Plasmid R702 is from Proteus mirabilis and can replicate in E.coli (Villarroel et al (1983) Mol. Gen. Genet. 189:390). Plasmid R702::pMc5BS obtained cointegration by through illegitimate recombination between pMc5BS and R702, mediated by transposable elements present on R702 (Leemans (1982) "Technieken voor het gebruik van Ti-plasmieden van Agrobacterium tumefaciens als vectoren voor genetic engineering van planten", Ph.D. Thesis Vrije Universiteit Brussel, Brussels, Belgium) and checked for induced expression of Barstar.

The use of E.coli (R702::pMc5BS) allows the construction, maintenance, amplification, and purification of plasmids containing the barnase DNA, such as pGSJVR1, without any lethal effect on the host due to accidental expression of the barnase DNA. However, because the Zml3 promoter is not expressed in E. coli, all steps of vector construction involving this promoter are also carried out in E. coli strain MC1061.

Example 3: Transformation of corn with the maintainer gene of Example 2.

Zygotic immature embryos of about 0.5 to 1 mm are isolated from developing seeds of corn inbred H99. The freshly isolated embryos enzymatically treated for 1-2 minutes with an enzyme solution II (0.3% macerozyme (Kinki Nishinomiya, Japan) in CPW salts (Powell & Chapman (1985) "Plant Cell Culture, A Practical Approach", R.A. Dixon ed., Chapter 3) with 10% mannitol and 5 mM 2-[N-Morpholino] ethane sulfonic acid (MES), pH 5.6).

After 1-2 minutes incubation in this enzyme solution, the embryos are carefully washed with N6aph solution (macro- and micro-elements of N6 medium (Chu et al (1975) Sci. Sin. Peking 18:659) supplemented with 6mM asparagine, 12 mM proline, 1 mg/l thiamine-HCl, 0.5 mg/l nicotinic acid, 100 mg/l casein hydrolysate, 100 mg/l inositol, 30 q/l sucrose and 54 g/l mannitol). After washing, the embryos are incubated in the maize electroporation buffer, EPM-KCl (80 mM KCl, 5 mM mM HEPES (N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid) and 0.425 M mannitol, pH 7.2). Approximately 100 embryos in 200 \(\mu \)l EPM-KCl are loaded in each electroporation cuvette. About 20 μg of a plasmid DNA, pPGSJVR1 (of Example 2) linearized with EcoRI, is added per cuvette.

After 1 hour DNA incubation with the explants, the cuvettes are transferred to an ice bath. After 10 minutes incubation on ice, the electroporation is carried out: one pulse with a field strength of 375 V/cm is discharged from a 900 μ F capacitor. The electroporation apparatus is as described by Dekeyser et al (1990) The Plant Cell 2:591. Immediately after electroporation, fresh liquid N6aph substrate is added to the explants in the cuvette, after which the explants are incubated for a further 10 minute period on ice.

Afterwards, the embryos are transferred to Mahl VII substrate (macro- and micro-elements and vitamins of N6 medium supplemented with 100 mg/l casein hydrolysate, 6 mM proline, 0.5 g/l MES, 1 mg/l 2,4-dichlorophenoxyacetic acid (2,4-D) and 2% sucrose solidified with 0.75 g/l MgCl₂ and 1.6 g/l Phytagel (Sigma Chemical Company, St Louis, Mo. U.S.A.), pH 5.8) and supplemented with 0.2M mannitol. After 3 days, the embryos are transferred to the same substrate supplemented with 200 mg/l kanamycin. After

approximately 14 days, the embryos are transferred to Mahl VII substrate without mannitol, supplemented with kanamycin. The embryos are further subcultured this selective substrate for approximately 2 months with subculturing intervals of about 3 weeks. The induced embryogenic tissue is carefully isolated and transferred to MS medium (Murashige and Skoog Physiol. Plant 15:473) supplemented with 5 (1962) mg/16-benzylaminopurine for line embryogenic tissue is maintained on this medium for approximately 14 days and subsequently transferred to MS medium without hormones and 6% sucrose for line H99. Developing shoots are transferred to 1/2 MS medium with 1.5% sucrose for further development to normal plantlets. These plantlets are transferred to soil and cultivated in the greenhouse.

In an analogous way, corn embryos are transformed with a fragment of pTS218 DNA which contains the maintainer gene and the chimeric P35S-bar-3'nos and which is obtained by digestion of the plasmid with EcoRI, XhoI and PstI and by purifying the longest fragment. Transformation and plant regeneration is as described in Example 5.

Example 4: Analysis of the transgenic corn plants of Example 3.

Plants form Example 3 transformed with pGSJVRl are analysed for the presence of the maintainer gene by means of PCR. DNA is prepared according to the protocol described by Dellaporta et al (1983) Plant Mol. Biol. Reporter 1:19, adapted for application to tissue amounts of about 10 to 20 mg. For each plant, such an amount of tissue is macerated in extraction buffer in a microfuge tube. Representative fragments of the maintainer gene are amplified using appropriate oligonucleotide probes.

Activity of the expression product of the first marker gene (i.e., neomycin phosphotransferase II (NPTII)) is assayed in plants as follows. Crude extracts are prepared by grinding plant tissue in extraction buffer (McDonnell et al (1987) Plant Molecular Biol. Reporter 5:380). The extracts are then subjected to non-denaturing polyacrylamide gel electrophoresis according to the procedure described by Reiss et al (1984) Gene 30:211. NPTII activity is then assayed by in situ phosphorylation of kanamycin using [gamma-32P]ATP as a substrate (McDonnell et al (1987) supra).

The plants that are found to be positive on both the PCR and NPTII assay are further analyzed by means

of Southern hybridization. Genomic DNA is prepared from plant tissue according to the protocol described by Dellaporta et al (1983) supra, supplemented by a treatment with RNase to remove remaining RNA. A nontransformed H99 plant is used as a control. Samples of the DNA are digested with appropriate restriction subjected to horizontal enzymes and electrophoresis. Southern transfer to Hybond N+ PLC, Amersham, International (Amersham Kingdom) membranes by means of the "alkali blotting of DNA" protocol and the subsequent hybridization are the manufacturer by recommended performed as (Amersham Hybond-N+ leaflet). Suitable radioactive prepared with the multi-prime are labelling kit (Amersham) according to the protocol supplied by the manufacturer which is derived from published procedures (Feinberg and Vogelstein (1983) Anal. Biochem. 132:6). The banding patterns show that at least the maintainer gene is integrated into the plant genomic DNA.

The PCR assays show that the maintainer gene is present. The NPTII assays show that the first marker DNA is expressed. The mature transformed plants can then be analyzed phenotypically to see whether the barstar DNA is expressed in tapetum cells and the barnase gene is expressed in pollen cells. Expression of barstar is determined by northern blotting of anther mRNA and by making testcrosses to determine the restoration in the progeny. Expression of the pollen-lethality gene is determined by cytological examination of the anther. In this regard, viable and nonviable mature pollen is determined by analyzing the staining of isolated pollen upon incubation for 30 minutes at 24°C in the following reaction mixture: 100 phosphate buffer Hq 7.8, 100 mM mM

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Sodiumsuccinate and 1 mm NitroBlue Tetrazolium, inspection by visual followed Alternative pollen. viable precipitation in techniques for the differentiation between viable and nonviable mature pollen are those described for example by Alexander (1969) Stain Technology 44:117, and by Heslop-Harrison and Heslop-Harrison (1970) Stain Technology 45:115. The viability of microspores is determined by embedding flower buds in plastic at different developmental stages and subjecting the buds to histochemical staining with the succinate dehydrogenase assay, both as described by De Block and Debrouwer (1992) The Plant Journal 2:261.

Ultimately, the progeny of the plant transformed with the pollen-lethality gene is determined. None of the offspring obtained from a cross using this plant as a male parent have this gene, while 50% of the offspring obtained from a cross using this plant as a female parent possess the gene.

Plants from Example 3, transformed with pTS218 DNA, are analyzed in the same way, except that the expression product of the first marker gene, i.e., phosphinothricine acetyltransferase, is assayed by means of a PAT assay as described in Example 5.

Example 5 : Production of male-sterile corn plants.

Zygotic embryos of corn inbred line H99 were isolated, enzymatically treated, washed, and loaded in electroporation buffer as described in Example 3. Approximately 100 embryos in 200 µl EPM-KCl were loaded in each electroporation cuvette. About 20 µg of a plasmid DNA, pVE108 linearized with HindIII, was added per cuvette. pVE108 is a 5620 bp plasmid which contains: a chimaeric gene comprising the bar DNA (EP 242236), encoding phosphinothricin acetyl transferase (PAT) and conferring resistance to an herbicidal inhibitor qlutamine synthetase such phosphinothricin (PPT), under the control of the 3553 promoter; and another chimaeric gene comprising the DNA coding for barnase (Hartley (1988) supra) under the control of the tapetum-specific promoter of the

TA29 gene (EP 344029) of N. tabacum. The complete sequence of plasmid pVE108 is given in SEQ ID no. 4. All vector constructions involving DNA fragments comprising the barnase gene were carried out in E. MC1061 containing the coli strain R702::pMc5BS of Example 3. After a 1 hour DNA incubation with the explants, the cuvettes were transferred to an ice bath. After 10 incubation on ice, the electroporation was carried out as described in Example 3. Immediately after electroporation, fresh liquid N6aph substrate was added to the explants in the cuvette, after which the explants were incubated for a further 10 minute period on ice.

Afterwards, the embryos from one electroporation experiment were transferred to Mahl VII substrate supplemented with 0.2 M mannitol and 2 mg/l PPT. After approximately 14 days, the embryos transferred to Mhl VII substrate (Mahl VII substrate Example 3 but without proline and hydrolysate) supplemented with 2 mg/l PPT but without mannitol. After approximately 4 weeks, the embryos subcultured for another month on Mh1 substrate supplemented with 10 mg/l PPT. The induced embryogenic tissue was carefully isolated transferred to MS medium supplemented with 5 mg/l 6benzylaminopurine. embryogenic The tissue maintained on this medium for approximately 14 days and subsequently transferred to MS medium without hormones and sucrose. Developing shoots transferred to 1/2 MS medium with 1.5% sucrose for development to normal plantlets. plantlets survived an in vitro spraying with doses of BASTAR (Hoechst AG, Frankfurt am Main, Germany) corresponding to 2 1/ha. These plantlets were then transferred to soil and cultivated in the greenhouse, and two of the transformed plantlets, designated RZM35-1 and RZM35-18, were further characterized.

embryos from a second electroporation experiment were transferred to Mhl VII substrate supplemented with 2 mg/l PPT and 0.2 M mannitol. After about 14 days, the embryos were transferred to Mh1 VII substrate supplemented with 2 mg/l PPT but without mannitol. After approximately another three weeks, the embryos were transferred to Mh1 substrate supplemented with 10 mg/l PPT but without mannitol. After another three weeks, the induced carefully isolated embryogenic tissue was transferred to MS medium supplemented with 2 mg/l PPT 5 mg/l 6-benzylaminopurine. The embryogenic and maintained on this medium approximately 14 days and subsequently transferred to without hormones, sucrose Developing shoots were transferred to 1/2 MS medium with 1.5% sucrose for further development to normal plantlets. The resulting plantlets were transferred to soil and cultivated in the greenhouse, and three of the transformed plantlets, designated RZM34-1, RZM34-12, and RZM34-14, were further characterized.

RZM34-1, RZM34-12, RZM34-14, RZM35-1, and RZM35-18 were grown in the greenhouse. Activity of the expression product of the <u>bar</u> gene in leaves of the plants was assayed as follows in a "PAT assay". 100 mg of leaf tissue from each plant, together with 50 mg of acid-treated sea sand (Merck, Darmstadt, Germany) and 5 mg polyvinylpolypyrrolidone (PVPP), were ground in an Eppendorf tube with a glass rod in 50 μ l of extraction buffer (25 mM Tris-HCL pH 7.5, 1 mM Na₂-EDTA (ethylenediaminetetraacetic acid disodium salt), 0.15 mg/ml phenylmethylsulfonylfluoride

(PMSF), 0.3 mg/ml dithiothreitol (DTT), and 0.3 mg/ml bovine serum albumin). The extract was centrifuged in microfuge for 5 minutes at 16000 rpm. supernatant was recovered and diluted ten times with TE 25/1 (25 mM Tris-HCL pH 7.5, 1 mM Na₂-EDTA. To twelve μ l of the diluted extract was then added: 1μ l of 1 mM PPT in TE 25/1, 1 μ l of 2 mM AcetylCoenzyme A in TE 25/1, and 2 μ l of [14C]AcetylCoenzym A (60 mCi/mmol, 0.02 mCi/ml, INEN Research Dupont, Wilmington, Delaware, USA). The reaction mixture was incubated for 30 minutes at 37°C and spotted on a aluminium sheet silicagel 60 t.l.c. plate with concentrating zone (Merck). Ascending chromatography was carried out in a 3 to 2 mixture of 1-propanol and NH₂OH (25% NH₇). 14C was visualized by overnight autoradiography (XAR-5 Kodak film). tolerance to the herbicide BASTAR was tested by brushing a small area near the top of one leaf per plant with a 1% solution of the herbicide and observing the damage symptoms at and near the brushed sites. While RZM34-1, RZM35-1 and RZM35-18 showed no damage symptoms at all, RZM34-12 and RZM34-14 displayed slight browning and drying-out of brushed site. RZM34-1, RZM34-12, RZM34-14, RZM35-1 and RZM35-18 were also shown to be male-sterile but otherwise phenotypically completely normal; female fertility, for instance, was normal. The spikelets of the male flowers were of about normal length but were very thin and appeared to be empty, and they never opened. A detailed analysis showed that the anthers were reduced to almost microscopic structures. This phenotype indicates not only that at least one copy of the barnase gene was expressed but also that it was selectively expressed in some or all of the tissues of the anthers.

Southern analysis showed RZM35-1 and RZM35-18 to have an identical integration pattern, with only one copy of plasmid pVE108 being present in the genome of each plant. A small part of the plasmid DNA sequence adjacent to the HindIII site (used for linearization prior to electroporation) seemed to be absent in the copy. Southern analysis of RZM34-1, integrated RZM34-12 and RZM34-14 showed that each of these plants probably has two or three copies of part or all of pVE108 integrated into its genome. The copies likely not inserted in а concatemer most configuration.

Transformants RZM35-1 and RZM34-1 were pollinated with pollen from an untransformed H99 plant, and progeny plantlets were recovered. From the 35 plantlets recovered from RZM35-1, 16 (46%) scored positive in a PAT assay, while 19 (54%) were PAT negative. This proportion in the F1 progeny does not differ significantly from the 1:1 ratio expected under normal Mendelian segregation of one active copy of the chimaeric bar gene (X2 = 0.26).

From the 34 plantlets recovered from RZM34-1, 19 (56%) scored positive in a PAT assay, while 15 (44%) were PAT negative. This proportion in the F1 progeny does not differ significantly from the 1:1 ratio expected under normal Mendelian segregation, assuming that the transformed female parent had one active copy, or alternatively multiple active, but closely linked copies, of the chimaeric bar gene (X2 = 0.47).

Example 6 : Production of restorer corn plants.

Zygotic embryos of corn inbred line H99 were isolated, enzymatically treated, washed and loaded in electroporation buffer as described in Example 5. Approximately 100 embryos in 200 μ l EPM-KCl were loaded in each electroporation cuvette. About 20 μ g

of a plasmid DNA, pVE144 linearized with HindIII, was added per cuvette. pVE144 is a 6555 bp plasmid which was described in Example 2.

The embryos were electroporated, and the transformed cells were selected, grown into callus, and regenerated as described in Example 3. Transgenic plants were analyzed for the presence of the fertility-restorer gene and the marker gene by means of Southern hybridization and PCR. The expression of the fertility-restorer gene is assayed by means of Northern blotting, and the expression of the marker gene is determined by NPTII assay as described in Example 3.

Example 7: Production of maintainer corn plants and a male-sterile corn line and maintenance of the male-sterile corn line

Maintainer plants of this invention of corn line H99 are obtained as outlined in Figure 1. A plant of corn inbred line H99 with the male-sterility genotype H99^{S/s,r/r,p/p}, transformed with the male-sterility gene is crossed with plants with the Example 5, H998/8,R/r,p/p, transformed with the genotype fertility-restorer gene of Example 6. The progeny that have the genotype H99^{S/s,R/r,p/p} are identified by PCR analysis for the presence of the S and R genes. These plants are selfed, yielding progeny with nine Two of these genotypes different genotypes. (H99^{S/S,r/r} and H99^{S/s,r/r}) will develop into male-sterile plants, while all the other genotypes will develop into male-fertile plants. When these male-fertile plants are selfed, progeny analysis allows identification of their genotype. Thus: a) progeny of selfings of $H99^{S/S,R/R}$, $H99^{S/s,R/R}$, $H99^{S/s,R/R}$, H998/8,R/r and H998/8,r/r would all develop into malefertile plants; b) selfings of H995/s,R/r plants would produce progeny, of which 13 out of 16 would be male-fertile, and since the male-sterility gene is linked to the herbicide resistance gene, <u>bar</u>, 4 out of the 13 male-fertile plants would be sensitive to the herbicide BASTA^R; and c) selfings of H99^{S/S,R/r} plants would produce progeny, of which 12 out 16 would be fertile (4 out of 16 would have the genotype H99^{S/S,R/R} and 8 out of 16 would have the genotype H99^{S/S,R/r}), all of which would be resistant to the herbicide, and the male-sterile progeny of which (4 out of 16) would all be homozygous for the male-sterility gene (H99^{S/S,r/r}).

The homozygous male-sterile progeny (H99^{S/S,r/r}) of selfing (c) are then crossed with their male-fertile siblings, and only when the cross is with plants with the genotype H99^{S/S,R/r} are the resulting plants 50% male-sterile (all with the genotype H99^{S/S,r/r}) and 50% male-fertile (all with the genotype H99^{S/S,R/r}. Indeed, the alternative cross between H99^{S/S,R/r} and H99^{S/S,R/R} would result in 100% male-fertile progeny plants.

Maintainer plants are selected by crossing the plant with the genotype H99^{S/s,R/r,p/p} with a plant that is heterozygous for the maintainer gene of Example 2, i.e., (H99^{s/s,r/r,P/p}), using the latter plant as the The offspring with the genotype female parent. H99S/s,r/r,P/p are selected by means of testcrosses supplemented with PCR analysis of the progeny (which and Southern can be easily identified by PCR blotting for the presence of the S and P genes and the absence of the R gene). The selected fertile offspring are then selfed. One out of eight offspring have the desired genotype for a maintainer plant of this invention (H99^{S/S,P/P)} and can be further selected by means of testcrosses and PCR analysis of the

progeny. Indeed, only plants with this genotype will produce 50% male-sterile offspring (all H99^{\$/\$,p/p}) and 50% male-fertile offspring (all H99^{\$/\$,p/p}), thus growing at once both the desired homozygous male-sterile line and the maintainer line of this invention. Testcrosses also include the pollination of wild type H99 plants with pollen of the progeny plants obtained from the selfing of H99^{\$/\$,p/p} plants.

Homozygous male-sterile plants with the genotype H99^{\$/\$,r/r,p/p} are then pollinated by maintainer plants (H99^{\$/\$,r/r,p/p}) of this invention. All progeny have the genotype H99^{\$/\$,r/r,p/p}, so that the male-sterile line is maintained, as desired.

Example 8: Introduction of the male-sterility gene and the maintainer gene in inbred corn lines through classical breeding.

The male-sterility gene of Example 5 and the maintainer gene of Example 2 are transferred from corn inbred line H99 to another corn inbred line (A) by repeated backcrossings as follows. The maintainer plant H99^{S/S,P/P} is crossed as a female parent with an untransformed plant of line A (As/s,p/p). The offspring the genotype A-H99^{S/s,P/p} are with selected screening, using PCR, for the presence of both the maintainer gene (P) and the male-sterility gene (S). These plants are then crossed again as female parents with A^{5/8,p/p} plants, and the offspring that heterozygous for both the P and S genes are again selected by PCR. This process of backcrossing is repeated until finally plants with the genotype A^{\$/s,P/p} are obtained. These plants are then selfed, and the progeny are analyzed in the same way as described in Example 7. In this way, male-sterile plants with the genotype $A^{S/S,p/p}$ and maintainer plants of this invention with the genotype $A^{S/S,P/p}$ are obtained.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: PLANT GENETIC SYSTEMS N.V.
 - (B) STREET: Jozef Plateaustraat 22
 - (C) CITY: Ghent
 - (E) COUNTRY: Belgium
 - (F) POSTAL CODE (ZIP): 9000
 - (G) TELEPHONE: 32 91 358411
 - (H) TELEFAX: 32 91 240694
 - (I) TELEX: 11.361 Pgsgen
- (ii) TITLE OF INVENTION: Maintenance of male-sterile plants
- (iii) NUMBER OF SEQUENCES: 17
- (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
- (vi) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 07/899,072
 - (B) FILING DATE: 12-JUN-1992
- (vi) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 07/970,849
 - (B) FILING DATE: 03-NOV-1992

(2) INFORMATION FOR SEO ID NO: 1:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2661 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Zea mays
 - (B) STRAIN: inbred line W-22
 - (x) PUBLICATION INFORMATION:
 - (A) AUTHORS: Hamilton et al.,
 - (C) JOURNAL: Sex Plant Reprod.
 - (D) VOLUME: 2
 - (F) PAGES: 208-
 - (G) DATE: 1989
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

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(2) INFORMATION FOR SEQ ID NO: 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6555 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: circular
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: plasmid pVE144 (replicable in E.coli)
 - (ix) FEATURE:
 - (A) NAME/KEY: -

 - (B) LOCATION: 1..396
 (D) OTHER INFORMATION: /label= pUC18 /note= "pUC18 derived sequence"
 - (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: complement (397..751)
 - (D) OTHER INFORMATION: /label= 3'nos /note= "3' regulatory sequence containing the polyadenylation site derived from Agrobacterium T-DNA nopaline synthase gene*
 - (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: complement (752..1024)
 - (D) OTHER INFORMATION: /label= barstar

/note= "coding region of the barstar gene of Bacillus amyloliquefaciens"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (1025..1607)
- (D) OTHER INFORMATION: /label= TA29

/note= "promoter derived from the TA29 gene of Nicotiana tabacum"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 1608..2440
- (D) OTHER INFORMATION: /label= 35S3 /note= *35S3 promoter sequence d

/note= "35S3 promoter sequence derived from cauliflower mosaic virus isolate CabbB-JI"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2441..3256
- (D) OTHER INFORMATION: /label= neo /note= "coding region of the neomycine phosphotransferase gene of Tn5"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 3257..4315
- (D) OTHER INFORMATION: /label= 3'ocs
 /note= "3' regulatory sequence containing the
 polyadenylation site derived from Agrobacterium
 T-DNA octopine synthase gene"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 4316..6555
- (D) OTHER INFORMATION: /label= pUC18 /note= "pUC18 derived sequence"

(xi) SEQUENCE DESCRIPTION: SEO ID NO: 2:

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CTATCTGTCA	CTTCATCGAA	AGGACAGTAG	AAAAGGAAGG	TGGCTCCTAC	AAATGCCATC	2160
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GCTCGACGTT	GTCACTGAAG	CGGGAAGGGA	CTGGCTGCTA	TTGGGCGAAG	TGCCGGGGCA	2760 ⁻
GGATCTCCTG	TCATCTCACC	TTGCTCCTGC	CGAGAAAGTA	TCCATCATGG	CTGATGCAAT	2820
GCGGCGGCTG	CATACGCTTG	ATCCGGCTAC	CTGCCCATTC	GACCACCAAG	CGAAACATCG	2880
CATCGAGCGA	GCACGTACTC	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	ATCTGGACGA	2940
AGAGCATCAG	GGGCTCGCGC	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	GCATGCCCGA	3000
CGGCGAGGAT	CTCGTCGTGA	CCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	3060
TGGCCGCTTT	TCTGGATTCA	TCGACTGTGG	CCGGCTGGGT	GTGGCGGACC	GCTATCAGGA	3120
CATAGCGTTG	GCTACCCGTG	ATATTGCTGA	AGAGCTTGGC	GGCGAATGGG	CTGACCGCTT	3180
CCTCGTGCTT	TACGGTATCG	CCGCTCCCGA	TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTCT	3240
TGACGAGTTC	TTCTGAGCGG	GACTCTGGGG	TTCGAAATGA	CCGACCAAGC	GACGCCCAAC	3300
CTGCCATCAC	GAGATTTCGA	TTCCACCGCC	GCCTTCTATG	AAAGGTTGGG	CTTCGGAATC	3360
GTTTTCCGGG	ACGCCGGCTG	GATGATCCTC	CAGCGCGGGG	ATCTCATGCT	GGAGTTCTTC	3420
GCCCACCCCC	TGCTTTAATG	AGATATGCGA	GACGCCTATG	ATCGCATGAT	ATTTGCTTTC	3480
AATTCTGTTG	TGCACGTTGT	AAAAAACCTG	AGCATGTGTA	GCTCAGATCC	TTACCGCCGG	3540
TTTCGGTTCA	TTCTAATGAA	TATATCACCC	GTTACTATCG	TATTTTTATG	AATAATATTC	3600
TCCGTTCAAT	TTACTGATTG	TACCCTACTA	CTTATATGTA	CAATATTAAA	ATGAAAACAA	3660
TATATTGTGC	TGAATAGGTT	TATAGCGACA	TCTATGATAG	AGCGCCACAA	TAACAAACAA	3720
TTGCGTTTTA	TTATTACAAA	TCCAATTTTA	AAAAAAGCGG	CAGAACCGGT	CAAACCTAAA	3780
AGACTGATTA	CATAAATCTT	ATTCAAATTT	CAAAAGGCCC	CAGGGGCTAG	TATCTACGAC	3840
ACACCGAGCG	GCGAACTAAT	AACGTTCACT	GAAGGGAACT	CCGGTTCCCC	GCCGGCGCGC	3900
ATGGGTGAGA	TTCCTTGAAG	TTGAGTATTG	GCCGTCCGCT	CTACCGAAAG	TTACGGGCAC	3960
CATTCAACCC	GGTCCAGCAC	GGCGGCCGGG	TAACCGACTT	GCTGCCCCGA	GAATTATGCA	4020
GCATTTTTTT	GGTGTATGTG	GGCCCCAAAT	GAAGTGCAGG	TCAAACCTTG	ACAGTGACGA	4080
CAAATCGTTG	GGCGGGTCCA	GGGCGAATTT	TGCGACAACA	TGTCGAGGCT	CAGCAGGGC	4140
TCGATCCCCT	CGCGAGTTGG	TTCAGCTGCT	GCCTGAGGCT	GGACGACCTC	GCGGAGTTCT	4200
ACCGGCAGTG	CAAATCCGTC	GGCATCCAGG	AAACCAGCAG	CGGCTATCCG	CGCATCCATG	4260
CCCCGAACT	GCAGGAGTGG	GGAGGCACGA	TGGCCGCTTT	GGTCGACCTG	CAGCCAAGCT	4320
TGGCGTAATC	ATGGTCATAG	CTGTTTCCTG	TGTGAAATTG	TTATCCGCTC	ACAATTCCAC	4380

ACAACATACG	AGCCGGAAGC	ATAAAGTGTA	AAGCCTGGGG	TGCCTAATGA	GTGAGCTAAC	4440
TCACATTAAT	TGCGTTGCGC	TCACTGCCCG	CTTTCCAGTC	GGGAAACCTG	TCGTGCCAGC	4500
TGCATTAATG	AATCGGCCAA	CGCGCGGGGA	GAGGCGGTTT	GCGTATTGGG	CGCTCTTCCG	4560
CTTCCTCGCT	CACTGACTCG	CTGCGCTCGG	TCGTTCGGCT	GCGGCGAGCG	GTATCAGCTC	4620
ACTCAAAGGC	GGTAATACGG	TTATCCACAG	AATCAGGGGA	TAACGCAGGA	AAGAACATGT	4680
GAGCAAAAGG	CCAGCAAAAG	GCCAGGAACC	GTAAAAAGGC	CGCGTTGCTG	GCGTTTTTCC	4740
ATAGGCTCCG	CCCCCTGAC	GAGCATCACA	AAAATCGACG	CTCAAGTCAG	AGGTGGCGAA	4800
ACCCGACAGG	ACTATAAAGA	TACCAGGCGT	TTCCCCCTGG	AAGCTCCCTC	GTGCGCTCTC	4860
CTGTTCCGAC	CCTGCCGCTT	ACCGGATACC	TGTCCGCCTT	TCTCCCTTCG	GGAAGCGTGG	4920
CGCTTTCTCA	ATGCTCACGC	TGTAGGTATC	TCAGTTCGGT	GTAGGTCGTT	CGCTCCAAGC	4980
TGGGCTGTGT	GCACGAACCC	CCCGTTCAGC	CCGACCGCTG	CGCCTTATCC	GGTAACTATC	5040
GTCTTGAGTC	CAACCCGGTA	AGACACGACT	TATCGCCACT	GGCAGCAGCC	ACTGGTAACA	5100
GGATTAGCAG	AGCGAGGTAT	GTAGGCGGTG	CTACAGAGTT	CTTGAAGTGG	TGGCCTAACT	5160
ACGGCTACAC	TAGAAGGACA	GTATTTGGTA	TCTGCGCTCT	GCTGAAGCCA	GTTACCTTCG	5220
GAAAAAGAGT	TGGTAGCTCT	TGATCCGGCA	AACAAACCAC	CGCTGGTAGC	GGTGGTTTTT	5280
TTGTTTGCAA	GCAGCAGATT	ACGCGCAGAA	AAAAAGGATC	TCAAGAAGAT	CCTTTGATCT	5340
TTTCTACGGG	GTCTGACGCT	CAGTGGAACG	AAAACTCACG	TTAAGGGATT	TTGGTCATGA	5400
GATTATCAAA	AAGGATCTTC	ACCTAGATCC	TTTTAAATTA	AAAATGAAGT	TTTAAATCAA	5460
TCTAAAGTAT	ATATGAGTAA	ACTTGGTCTG	ACAGTTACCA	ATGCTTAATC	AGTGAGGCAC	5520
CTATCTCAGC	GATCTGTCTA	TTTCGTTCAT	CCATAGTTGC	CTGACTCCCC	GTCGTGTAGA	5580
TAACTACGAT	ACGGGAGGGC	TTACCATCTG	GCCCCAGTGC	TGCAATGATA	CCGCGAGACC	5640
CACGCTCACC	GGCTCCAGAT	TTATCAGCAA	TAAACCAGCC	AGCCGGAAGG	GCCGAGCGCA	5700
GAAGTGGTCC	TGCAACTTTA	TCCGCCTCCA	TCCAGTCTAT	TAATTGTTGC	CGGGAAGCTA	5760
GAGTAAGTAG	TTCGCCAGTT	AATAGTTTGC	GCAACGTTGT	TGCCATTGCT	ACAGGCATCG	5820
TGGTGTCACG	CTCGTCGTTT	GGTATGGCTT	CATTCAGCTC	CGGTTCCCAA	CGATCAAGGC	5880
GAGTTACATG	ATCCCCCATG	TTGTGCAAAA	AAGCGGTTAG	CTCCTTCGGT	CCTCCGATCG	5940
TTGTCAGAAG	TAAGTTGGCC	GCAGTGTTAT	CACTCATGGT	TATGGCAGCA	CTGCATAATT	6000
CTCTTACTGT	CATGCCATCC	GTAAGATGCT	TTTCTGTGAC	TGGTGAGTAC	TCAACCAAGT	6060
CATTCTGAGA	ATAGTGTATG	CGGCGACCGA	GTTGCTCTTG	CCCGGCGTCA	ATACGGGATA	6120
ATACCGCGCC	ACATAGCAGA	ACTTTAAAAG	TGCTCATCAT	TGGAAAACGT	TCTTCGGGGC	6180
GAAAACTCTC	AAGGATCTTA	CCGCTGTTGA	GATCCAGTTC	GATGTAACCC	ACTCGTGCAC	6240

CCAACTGATC	TTCAGCATCT	TTTACTTTCA	CCAGCGTTTC	TGGGTGAGCA	AAAACAGGAA	6300
GGCAAAATGC	CGCAAAAAAG	GGAATAAGGG	CGACACGGAA	ATGTTGAATA	CTCATACTCT	6360
TCCTTTTTCA	ATATTATTGA	AGCATTTATC	AGGGTTATTG	TCTCATGAGC	GGATACATAT	6420
TTGAATGTAT	TTAGÁAAAAT	AAACAAATAG	GGGTTCCGCG	CACATTTCCC	CGAAAAGTGC	6480
CACCTGACGT	CTAAGAAACC	ATTATTATCA	TGACATTAAC	СТАТАААААТ	AGGCGTATCA	6540
CGAGGCCCTT	TCGTC					6555

(2) INFORMATION FOR SEQ ID NO: 3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5620 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: circular
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: plasmid pVE108 (replicable in E.coli)
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..395
 - (D) OTHER INFORMATION: /label= pUC18 /note= *pUC18 derived sequence*
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: complement (396..802)
 - (D) OTHER INFORMATION: /label= 3'nos /note= "3' regulatory sequence containing the polyadenylation site derived from the nopaline synthase gene from Agrobacterium T-DNA"
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: complement (803..1138)
 - (D) OTHER INFORMATION: /label= barnase /note= "coding region of the barnase gene of Bacillus amyloliquefaciens"
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: complement (1139..1683)
 - (D) OTHER INFORMATION: /label= TA29

/note= "sequence derived from tapetum specific
promoter of Nicotiana tabacum"

- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1684..2516
 - (D) OTHER INFORMATION: /label= 35S3

/note= ""35S3" promoter sequence derived from cauliflower mosaic virus isolate CabbB-JI"

(ix) FEATURE:

- (A) NAME/KEY: (B) LOCATION: 2517..3068
 (D) OTHER INFORMATION: /label= bar /note= "coding sequence of phosphinotricin acetyltransferase gene*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 3069..3356
- (D) OTHER INFORMATION: /label= 3'nos /note= "3' regulatory sequence containing the polyadenylation site derived from Agrobacterium T-DNA nopaline synthase gene"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 3357..5620
- (D) OTHER INFORMATION: /label= puc18 /note= "pUC18 derived sequence"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

TCGCGCGTTT	CGGTGATGAC	GGTGAAAACC	TCTGACACAT	GCAGCTCCCG	GAGACGGTCA	60
CAGCTTGTCT	GTAAGCGGAT	GCCGGGAGCA	GACAAGCCCG	TCAGGGCGCG	TCAGCGGGTG	120
TTGGCGGGTG	TCGGGGCTGG	CTTAACTATG	CGGCATCAGA	GCAGATTGTA	CTGAGAGTGC	180
ACCATATGCG	GTGTGAAATA	CCGCACAGAT	GCGTAAGGAG	AAAATACCGC	ATCAGGCGCC	240
ATTCGCCATT	CAGGCTGCGC	AACTGTTGGG	AAGGGCGATC	GGTGCGGGCC	TCTTCGCTAT	300
TACGCCAGCT	GGCGAAAGGG	GGATGTGCTG	CAAGGCGATT	AAGTTGGGTA	ACGCCAGGGT	360
TTTCCCAGTC	ACGACGTTGT	AAAACGACGG	CCAGTGAATT	CGAGCTCGGT	ACCCGGGGAT	420
CTTCCCGATC	TAGTAACATA	GATGACACCG	CGCGCGATAA	TTTATCCTAG	TTTGCGCGCT	480
ATATTTTGTT	TTCTATCGCG	TATTAAATGT	ATAATTGCGG	GACTCTAATC	ATAAAAACCC	540
ATCTCATAAA	TAACGTCATG	CATTACATGT	TAATTATTAC	ATGCTTAACG	TAATTCAACA	600
GAAATTATAT	GATAATCATC	GCAAGACCGG	CAACAGGATT	CAATCTTAAG	AAACTTTATT	660
GCCAAATGTT	TGAACGATCT	GCTTCGGATC	CTCTAGAGNN	NNCCGGAAAG	TGAAATTGAC	720
CGATCAGAGT	TTGAAGAAAA	ATTTATTACA	CACTTTATGT	AAAGCTGAAA	AAAACGGCCT	780
CCGCAGGAAG	CCGTTTTTTT	CGTTATCTGA	TTTTTGTAAA	GGTCTGATAA	TGGTCCGTTG	840
TTTTGTAAAT	CAGCCAGTCG	CTTGAGTAAA	GAATCCGGTC	TGAATTTCTG	AAGCCTGATG	900
TATAGTTAAT	ATCCGCTTCA	CGCCATGTTC	GTCCGCTTTT	GCCCGGGAGT	TTGCCTTCCC	960
TGTTTGAGAA	GATGTCTCCG	CCGATGCTTT	TCCCCGGAGC	GACGTCTGCA	AGGTTCCCTT	1020
TTGATGCCAC	CCAGCCGAGG	GCTTGTGCTT	CTGATTTTGT	AATGTAATTA	TCAGGTAGCT	1080

TATGATATGI	CTGAAGATAA	TCCGCAACCC	CGTCAAACGT	GTTGATAACC	GGTACCATGG	1140
TAGCTAATTI	CTTTAAGTAA	AAACTTTGAT	TTGAGTGATG	ATGTTGTACT	GTTACACTTG	1200
CACCACAAGG	GCATATATAG	AGCACAAGAC	ATACACAACA	ACTTGCAAAA	CTAACTTTTG	1260
TTGGAGCATT	TCGAGGAAAA	TGGGGAGTAG	CAGGCTAATC	TGAGGGTAAC	ATTAAGGTTT	1320
CATGTATTAA	TTTGTTGCAA	ACATGGACTT	AGTGTGAGGA	AAAAGTACCA	AAATTTTGTC	1380
TCACCCTGAT	TTCAGTTATG	GAAATTACAT	TATGAAGCTG	TGCTAGAGAA	GATGTTTATT	1440
CTAGTCCAGC	CACCCACCTT	ATGCAAGTCT	GCTTTTAGCT	TGATTCAAAA	ACTGATTTAA	1500
TTTACATTGC	TAAATGTGCA	TACTTCGAGC	CTATGTCGCT	TTAATTCGAG	TAGGATGTAT	1560
ATATTAGTAC	АТААААААТС	ATGTTTGAAT	CATCTTTCAT	AAAGTGACAA	GTCAATTGTC	1620
CCTTCTTGTT	TGGCACTATA	TTCAATCTGT	TAATGCAAAT	TATCCAGTTA	TACTTAGCTA	1680
GATCCTACGC	AGCAGGTCTC	ATCAAGACGA	TCTACCCGAG	TAACAATCTC	CAGGAGATCA	1740
AATACCTTCC	ÇAAGAAGGTT	AAAGATGCAG	TCAAAAGATT	CAGGACTAAT	TGCATCAAGA	1800
ACACAGAGAA	AGACATATTT	CTCAAGATCA	GAAGTACTAT	TCCAGTATGG	ACGATTCAAG	1860
GCTTGCTTCA	TAAACCAAGG	CAAGTAATAG	AGATTGGAGT	CTCTAAAAAG	GTAGTTCCTA	1920
CTGAATCTAA	GGCCATGCAT	GGAGTCTAAG	ATTCAAATCG	AGGATCTAAC	AGAACTCGCC	1980
GTGAAGACTG	GCGAACAGTT	CATACAGAGT	CTTTTACGAC	TCAATGACAA	GAAGAAAATC	2040
TTCGTCAACA	TGGTGGAGCA	CGACACTCTG	GTCTACTCCA	AAAATGTCAA	AGATACAGTC	2100
TCAGAAGACC	AAAGGGCTAT	TGAGACTTTT	CAACAAAGGA	TAATTTCGGG	AAACCTCCTC	2160
GGATTCCATT	GCCCAGCTAT	CTGTCACTTC	ATCGAAAGGA	CAGTAGAAAA	GGAAGGTGGC	2220
TCCTACAAAT	GCCATCATTG	CGATAAAGGA	AAGGCTATCA	TTCAAGATGC	CTCTGCCGAC	2280
AGTGGTCCCA	AAGATGGACC	CCCACCCACG	AGGAGCATCG	TGGAAAAAGA	AGACGTTCCA	2340
ACCACGTCTT	CAAAGCAAGT	GGATTGATGT	GACATCTCCA	CTGACGTAAG	GGATGACGCA	2400
CAATCCCACT	ATCCTTCGCA	AGACCCTTCC	TCTATATAAG	GAAGTTCATT	TCATTTGGAG	2460
AGGACACGCT	GAAATCACCA	GTCTCTCTCT	ATAAATCTAT	CTCTCTCTCT	ATAACCATGG	2520
ACCCAGAACG	ACGCCCGGCC	GACATCCGCC	GTGCCACCGA	GGCGGACATG	CCGGCGGTCT	2580
GCACCATCGT	CAACCACTAC	ATCGAGACAA	GCACGGTCAA	CTTCCGTACC	GAGCCGCAGG	2640
AACCGCAGGA	GTGGACGGAC	GACCTCGTCC	GTCTGCGGGA	GCGCTATCCC	TGGCTCGTCG	2700
CCGAGGTGGA	CGGCGAGGTC	GCCGGCATCG	CCTACGCGGG	CCCCTGGAAG	GCACGCAACG	2760
CCTACGACTG	GÀCGGCCGAG	TCGACCGTGT	ACGTCTCCCC	CCGCCACCAG	CGGACGGGAC	2820
TGGGCTCCAC	GCTCTACACC	CACCTGCTGA	AGTCCCTGGA	GGCACAGGGC	TTCAAGAGCG	2880
TGGTCGCTGT	CATCGGGCTG	CCCAACGACC	CGAGCGTGCG	CATGCACGAG	GCGCTCGGAT	2940

ATGCCCCCCG	CGGCATGCTG	CGGGCGGCCG	GCTTCAAGCA	CGGGAACTGG	CATGACGTGG	3000
GTTTCTGGCA	GCTGGACTTC	AGCCTGCCGG	TACCGCCCCG	TCCGGTCCTG	CCCGTCACCG	3060
AGATCTGATC	TCACGCGTCT	AGGATCCGAA	GCAGATCGTT	CAAACATTTG	GCAATAAAGT	3120
TTCTTAAGAT	TGAATCCTGT	TGCCGGTCTT	GCGATGATTA	TCATATAATT	TCTGTTGÄAT	3180
TACGTTAAGC	ATGTAATAAT	TAACATGTAA	TGCATGACGT	TATTTATGAG	ATGGGTTTTT	3240
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AACTAGGATA	AATTATCGCG	CGCGGTGTCA	TCTATGTTAC	TAGATCGGGA	AGATCCTCTA	3360
GAGTCGACCT	GCAGGCATGC	AAGCTTGGCG	TAATCATGGT	CATAGCTGTT	TCCTGTGTGA	3420
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TGGGGTGCCT	AATGAGTGAG	CTAACTCACA	TTAATTGCGT	TGCGCTCACT	GCCCGCTTTC	3540
CAGTCGGGAA	ACCTGTCGTG	CCAGCTGCAT	TAATGAATCG	GCCAACGCGC	GGGGAGAGGC	3600
GGTTTGCGTA	TTGGGCGCTC	TTCCGCTTCC	TCGCTCACTG	ACTCGCTGCG	CTCGGTCGTT	3660
CGGCTGCGGC	GAGCGGTATC	AGCTCACTCA	AAGGCGGTAA	TACGGTTATC	CACAGAATCA	3720
GGGGATAACG	CAGGAAAGAA	CATGTGAGCA	AAAGGCCAGC	AAAAGGCCAG	GAACCGTAAA	3780
AAGGCCGCGT	TGCTGGCGTT	TTTCCATAGG	CTCCGCCCCC	CTGACGAGCA	TCACAAAAAT	3840
CGACGCTCAA	GTCAGAGGTG	GCGAAACCCG	ACAGGACTAT	AAAGATACCA	GGCGTTTCCC	3900
CCTGGAAGCT	CCCTCGTGCG	CTCTCCTGTT	CCGACCCTGC	CGCTTACCGG	ATACCTGTCC	3960
GCCTTTCTCC	CTTCGGGAAG	CGTGGCGCTT	TCTCAATGCT	CACGCTGTAG	GTATCTCAGT	4020
TCGGTGTAGG	TCGTTCGCTC	CAAGCTGGGC	TGTGTGCACG	AACCCCCCGT	TCAGCCCGAC	4080
CGCTGCGCCT	TATCCGGTAA	CTATCGTCTT	GAGTCCAACC	CGGTAAGACA	CGACTTATCG	4140
CCACTGGCAG	CAGCCACTGG	TAACAGGATT	AGCAGAGCGA	GGTATGTAGG	CGGTGCTACA	4200
GAGTTCTTGA	AGTGGTGGCC	TAACTACGGC	TACACTAGAA	GGACAGTATT	TGGTATCTGC	4260
GCTCTGCTGA	AGCCAGTTAC	CTTCGGAAAA	AGAGTTGGTA	GCTCTTGATC	CGGCAAACAA	4320
ACCACCGCTG	GTAGCGGTGG	TTTTTTTGTT	TGCAAGCAGC	AGATTACGCG	CAGAAAAAA	4380
GGATCTCAAG	AAGATCCTTT	GATCTTTTCT	ACGGGGTCTG	ACGCTCAGTG	GAACGAAAAC	4440
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					GTCTGACAGT	4560
					TTCATCCATA	4620
	TCCCCGTCGT					4680
					AGCAATAAAC	4740
CAGCCAGCCG	GAAGGGCCGA	GCGCAGAAGT	GGTCCTGCAA	CTTTATCCGC	CTCCATCCAG	4800

TCTATTAATT	GTTGCCGGGA	AGCTAGAGTA	AGTAGTTCGC	CAGTTAATAG	TTTGCGCAAC	4860
GTTGTTGCCA	TTGCTACAGG	CATCGTGGTG	TCACGCTCGT	CGTTTGGTAT	GGCTTCATTC	4920
AGCTCCGGTT	CCCAACGATC	AAGGCGAGTT	ACATGATCCC	CCATGTTGTG	CAAAAAAGCG	4980
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ATGGTTATGG	CAGCACTGCA	TAATTCTCTT	ACTGTCATGC	CATCCGTAAG	ATGCTTTTCT	5100
GTGACTGGTG	AGTACTCAAC	CAAGTCATTC	TGAGAATAGT	GTATGCGGCG	ACCGAGTTGC	5160
TCTTGCCCGG	CGTCAATACG	GGATAATACC	GCGCCACATA	GCAGAACTTT	AAAAGTGCTC	5220
ATCATTGGAA	AACGTTCTTC	GGGGCGAAAA	CTCTCAAGGA	TCTTACCGCT	GTTGAGATCC	5280
AGTTCGATGT	AACCCACTCG	TGCACCCAAC	TGATCTTCAG	CATCTTTTAC	TTTCACCAGC	5340
GTTTCTGGGT	GAGCAAAAAC	AGGAAGGCAA	AATGCCGCAA	aaaagggaat	AAGGCCGACA	5400
CGGAAATGTT	GAATACTCAT	ACTCTTCCTT	TTTCAATATT	ATTGAAGCAT	TTATCAGGGT	5460
TATTGTCTCA	TGAGCGGATA	CATATTTGAA	TGTATTTAGA	AAAATAAACA	AATAGGGGTT	5520
CCGCGCACAT	TTCCCCGAAA	AGTGCCACCT	GACGTCTAAG	AAACCATTAT	TATCATGACA	5580
TTAACCTATA	AAAATAGGCG	TATCACGAGG	CCCTTTCGTC			5620

(2) INFORMATION FOR SEQ ID NO: 4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 21 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: oligonucleotide MDB80
- (ix) FEATURE:

 - (A) NAME/KEY: -(B) LOCATION: 1..21
 - (D) OTHER INFORMATION: /label= MDB80 /note= "oligonucleotide designated as MDB80"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

CCGCTTGTCA GTGAATGTTG C

21

(2) INFORMATION FOR SEQ ID NO: 5:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs

WO 93/25695

	59	
·	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iii)	ANTI-SENSE: NO	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide MDB81	
(ix)	FEATURE: (A) NAME/KEY: - (B) LOCATION: 121 (D) OTHER INFORMATION: /label= MDB81 /note= "oligonucleotide designated as MDB81"	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
CCGAGGCC	AT GGTTGCCGCC G	21
(2) <u>INFO</u>	RMATION FOR SEQ ID NO: 6:	
	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iii)	ANTI-SENSE: NO	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide MDB82	
(ix)	FEATURE: (A) NAME/KEY: - (B) LOCATION: 121 (D) OTHER INFORMATION: /label= MDB82 /note= "oligonucleotide designated as MDB82"	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
ACGCATAGO	GC ATAGGATGAC G	21
(2) TNEOS	RMATION FOR SEQ ID NO: 7:	
	SEQUENCE CHARACTERISTICS:	

(i)

(A) LENGTH: 3627 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(iii)	HYPOTHETICAL:	NO
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(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Oryza sativa
- (B) STRAIN: Akihikari

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 1..2845
 (D) OTHER INFORMATION: /label= PT72 /note= *sequence comprising anther specific promoter PT72*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2733..2739
- (D) OTHER INFORMATION: /label= TATA /note= "TATA Box"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2765
- (D) OTHER INFORMATION: /note= "transcription initiation determined by primer extension*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2846..2848
- (D) OTHER INFORMATION: /label= ATG /note= "ATG start of translation of rice T72 gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

GACAATACAT	CAAGTAAATC	AAACATTACA	AATCAGAACC	TGTCTAAGAA	TCCATCTTAA	60
TTCAGAAAAA	AACTCAGATT	AGATGTTCAT	GCTTCCACCA	GAAGCAGGAA	TGTGCAACCT	120
ACACTTCCTG	TAATTTCCAT	ACTACAATGT	CCCCACTGAC	CACTGTGCCT	GATGCTCTAT	180
TAGAATACCA	CATCCTCCAT	GGCTCCATGT	AAATGCATAT	AAATTTGACT	CTTTAAATTA	240
GTAACTACAA	TTTAAAATTT	ATCGAACATT	GTTCAAATTT	ATAAACAGTT	TCCCCAAATT	300
TAGATGCTCC	CAAATGTACA	CAGCTACTAG	TAAAGCACCA	TCCAGTTTCA	CCTGAACAGG	360
ACTGACATAA	ATGTGTGAAA	AGGGGACGTC	ATTCCCCCAA	ATACAACTGA	ACAATCCTCC	420
ATCAGAACAT	TCATTTGATT	GACATTACTC	GGAGAGATAC	AGCTCGCAGG	CACACGAGAT	480
TCTTCTGCCT	TTCCAATTGC	CACGAACCCA	CATGTCACAC	GACCAACCAA	AAAGAGAGAA	540
TTTTTCTTTG	CACAAAÇAAA	AAGTGAGATT	TTTTTTTCGC	CACAAAGGTG	CGAACTTTCT	600
TCTCTCTCCC	ACTTTCCAAT	CAAGAAACGA	AGCACTCAAA	CCAAGAACAA	ACCAAGGAAG	660
GAGAGATCGC	TCCCTCTCCC	AGAGCAAACG	AAAGGAGAGA	ACTCAGATGG	ATGCGAACTA	720
CTACCTTGCC	TCTTTCCCCG	GAGAAGCAGC	GAAGGAGAAG	AGCGCGATGC	CGCCGCCGCC	780
GCCGCCTCCG	GCAACCTCCG	GCTCCGGCGA	GTCCGCCTCC	TCCTCCTCTC	TCACCTCTCT	840

CTTCCCAACC	GTGTGGTGTT	CGAGAAGCTT	TTATGCGAGC	GACGTGCAGT	GGAAGCGGTT	900
GCTCCCAAGT	CAAACTGATG	GAGACCACCT	ACTATCTTCC	TCTTGTTTTC	TTCTGCTTTT	960
CTTTTCTTTA	TCTTTTTTCT	TTCATTTTAT	TTTGAGCGAT	GAACTTGAGA	ACAGTTTGGT	1020
TGTGGGTTAA	ATTAAACGGT	GCAGAATTGC	AAAGCTACGT	CCTTTTCGTC	TGATTAAGGT	1080
GGTATCAGAA	TCCTAATCTG	TTAGCTCAGC	ATTTGTTTTT	GTGTGTTTAA	TTGGCCATGA	1140
CATCAGATGG	TTCAGACCGG	TGGCAGGTCT	TCATCGGAGA	GGAGAATGAG	AGCAATGCAA	1200
GTTGCAAACA	ACAAACAGGT	CCTTCCAAAC	GGGTTGGTTT	CATTCCACAG	AACAGGATAG	1260
CAACCAGAGC	ACAAACCGTT	CAACAATATA	ТАТАТАТАТА	ТАТАТАТАТА	TATATATATA	1320
татататата	TATATATATG	ATTTAAAATT	ATATTACTAT	TTTTAGGATA	CGGAACTCTT	1380
AACACATGAA	AATCTAAACA	TTTTCAACCA	ATCAGAACTA	CTAGAAAGAT	AATCTAACTA	1440
CTTCAAAATT	TAAAATTTGA	CAAATAAAAT	AACTAGTTTT	TTCTAAAGCT	ATCTTCACTG	1500
GACAACTTAT	GAATATTTAT	ATTTATGAAG	CGAGTACTCT	CCTAGTACAT	ATTACATATA	1560
TATTCTTCTT	CTCATGAAAA	ATTAACTTCT	CGCTATAAAT	CCGAACATAT	ATTATGCGTA	1620
GCAAGTTGTT	TTTTTTAACG	GGTGGAGTAA	TATTAGAGTA	TTTAAATTCC	TTCAAATTGC	1680
CATCCCTCTG	GGACTTTGCT	GCTGTTGTTC	TTCCACGGTT	GCTGTCAGTG	TCACCCAGAT	1740
TTGCATCCTT	TCCAGCTCGT	AGCTACTGTT	CTGCATGTAT	TGGACTTGGA	TTAAGATCAA	1800
ATGCAGTTGC	TATTGTAACT	GCACAATAGC	AACTGCACAC	AATCATGTCC	ATTCGTTTTC	1860
AGATCCAACG	GCTCTAGATG	ACTGCTACAG	TACATGCATA	ATAGTACATC	TCTGCTACAG	1920
TGTTTTTGCT	GCAGTACCAC	TTCATATCCT	GCCTTCCGT	TCTAGATCAT	GTGATGTACA	1980
TGTTTTTTTG	AAACAACCCG	CACAAGACAT	TGATAGAGTA	GGAAATGTGA	TGTACATGTT	2040
AACGGCTTAA	GTTACAGTTA	CAATAACAAC	TGCACAGGAT	CTTGATCCAT	TGGACTTGTA	2100
TAATATCTCA	TCTCGTCGTT	CCATTATCGT	GGTAACAGTT	GGCAACTTGG	CATCCAGTGC	2160
TGGAAACTAT	GCCGTGTGTA	CATCAGGATC	GTCCTTTTTG	TTCAGTTCCA	AGATAGAACA	2220
AGTCCAAAAG	ATGGCCGTAG	TTTTTTTAGT	CACAGTGGAA	GCTGACATAG	CCGTGGAATA	2280
AGTTCTGCAC	AAAAGTTGCC	ATTCGAGATC	AACTACTGGT	AGTAGTAGTC	ATCTTCTACC	2340
ACTGCGAATA	TTCGAAGGGA	CACAAAAAGA	TCAACGAGTA	AATTAGTTCA	CCGGAAGACG	2400
ACACATTATC	ACCACAAAAA	GACTAAAAAC	AAAAAGAAAT	TGCCAGGCCA	AAAAAGGCAA	2460
AAAAGAAAAA	AAAAGATGGC	ACGAGGCCCA	GGGCTACGGC	CCATCTTGTC	GCCGGCCCAA	2520
CCGCGCGCGC	GAAACGCTCT	CGTCGGCTCT	CGGCTCGCCG	CGACGCGATG	GAGAGTTCGC	2580
GCCGCGGCGC	GCGCGCGCGT	TCGGTGGCTC	ACACGCTTGC	GCCCTCGTCC	TCCCGGCCGG	2640
CGCGGGCGCC	GACCGCGCGT	CCGCCGCATG	CGCGCGGCGT	AGGTGAGCAA	CGCGGGCCTC	2700

GCCGCGCGCG	CTCCCCTCCT	TCGATCCCCT	CCTATAAATC	GAGCTCGCGT	CGCGTATCGC	2760
CACCACCACC	ACGACACACA	CGCACGCACC	GTGCAGGCAT	CGACGACGAG	CGAGAGCCCC	2820
TCGGCGGCAG	AAGACACTCA	CGGCGATGGC	GGTGACGAGG	ACGGCGCTGC	TGGTGGTGTT	2880
GGTAGCGGGG	GCGATGACGA	TGACGATGCG	CGGGGCGGAG	GCGCAGCAGC	CGAGCTGCGC	2940
GGCGCAGCTC	ACGCAGCTGG	CGCCGTGCGC	GCGAGTCGGC	GTGGCGCCGG	CGCCGGGGCA	3000
GCCGCTGCCG	GCGCCCCCGG	CGGAGTGCTG	CTCGGCGCTG	GGCGCCGTGT	CGCACGACTG	3060
CGCCTGCGGC	ACGCTCGACA	TCATCAACAG	CCTCCCCGCC	AAGTGCGGCC	TCCCGCGCGT	3120
CACCTGCCGT	AAGAAAACGA	ATAAAATCGA	TTTGCTATCT	ATCGATGATT	GTGTTTTTGT	3180
AGACTAAACT	AAACCCCTAT	TAATAATCAA	CTAACCGATG	AACTGATCGT	TGCAGAGTGA	3240
TGGAGATGGT	GTGCCAAGGT	AATTGCGTTT	GCTCGTGCGA	GGATGAGAAG	AGAAGATTGA	3300
ATAAGATGTT	TGATGGCAAC	AAGTCATCAG	GCGATCCGAT	CCCTGCAGCT	ATGAATGGGA	3360
GTATACGTAG	TAGTGGTCTC	GTTAGCATCT	GTGTGTCGCA	TATGCACGCC	GTGCGTGCCG	3420
TGTCTGTCCT	GCTTGCTCTG	CTGATCGTTC	AATGAACGAC	AAATTAATCT	AACTCTGGAG	3480
TGACAAGTCG	TTCGAGATAT	ACTAATACTA	CCATGTGCAG	GGTCTTTCAA	CCAAGGTTCA	3540
TGTTTTCCAC	GAAAGCCGAT	TGAAACGAAA	CCGCGAAATT	TTGATGCGAG	ATGAAAGCAG	3600
ATTCCGAGTG	AAATTTTAAA	TGGTTTT				3627

(2) INFORMATION FOR SEQ ID NO: 8:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 2370 base pairs

 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Oryza sativa (B) STRAIN: Akihikari
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..1808
 - (D) OTHER INFORMATION: /label= PT42 /note= *sequence comprising anther specific promoter PT42*
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1748..1755
 - (D) OTHER INFORMATION: /label= TATA

/note= "TATA Box"

(ix) FEATURE:

- (A) NAME/KEY: -(B) LOCATION: 1780
- (D) OTHER INFORMATION: /note= "transcription initiation site determined by primer extension"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 1809
- (D) OTHER INFORMATION: /label= ATG

/note= "ATG start of translation of rice T42 gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

				4		
60	GCCGCCGTCG	TCCTTCCTGC	CATCACCGTC	GCGCCATGGA	GTCGGGTGCT	GGCCATCACT
120	TCTTCCCCAA	ACTACCTCTC	CCCTCACGCC	AGCCTTCTTC	CCAAGGCCGA	CCGGTGAGCT
180	AGCCCCGTGC	GCCATCCCAG	CCACCTCCAC	TGCCACAGCG	CGCCGTCCGT	TTCCGGCCAA
240	GCGGCAGCCA	GACGCTCGTC	TGCCAACGCC	CATCTCCTCT	GGTTCGCCTC	CGTGCCACCG
300	CCAGGGAGCC	CAGAGCACGG	ACAGCCATGG	CCGCCGCGCC	CCGATGAACA	TGCGCTGTCA
360	CTAGTCGGCT	GCAGCCGGAC	ACATCTGGTT	TCCTTCTCTC	TGCCTCCTCC	ATGGCTGCTC
420	CGTTTCAGTC	TTTCTCTCAC	TTTATGAAAG	AAAATTGTCT	GCCCATGGGC	TATACAAATG
480	TTTCGGTGTC	TACCATATTT	GCCAGCAAAT	AGGATTGTCC	AAATAATGGG	GGAAATAATA
540	CTAAGTGTCC	ACCACAATTT	CACAGCAAAG	TCGGGTGTTT	TACACGATCT	CAAGAGCAAA
600	ATGAAGAAAA	AAGACTGTAC	ACCAAAGGAG	AAATTTAAA	TTTGCCAATA	TGTAACAAAT
660	TTTAGGATGA	TTGATTTATT	GGTTATAAAG	TAAGCTCAGG	TGAAATTACA	ACAAAGAGAA
720	CTTGCTAAAA	ATATAACGTG	GTGTCGGAAA	GGCCAATTGG	TGAAAACAAT	AGGAAGTGTG
780	ATGCCCTGTA	GATGGTCAAG	GATAGACCCT	GCTGATTATA	ATATCCTGTA	TGTCGTCCCC
840	ACTCACATAT	TACTCCCCGA	TTCTCTCAAG	TCATCTCCGC	TTTCCATGCT	CTGGATCGTG
900	AGATAACCAA	CTCACTTCAT	TCAAACAACA	TAAGAAACAG	GGATCCACAG	CTGGTGGGCT
960	AAACTTGGTA	AAGTGCTTAA	TACTCCTAGT	CCTTATCTTA	ATTCTTAGTC	TTGTTTAATT
1020	GTAATTTTTA	CGTATACCAT	ATAATTATAA	CAATTACAAT	ATTTATCGTA	TAAATATCAA
1080	ACGCCGAACC	GCAGCAGCGG	ATGAGCAGCC	AAATATGGTG	TAGATAAAA	AAACTATTTT
1140	TTTGCTGGGT	TTAGTGTTCG	CTAAAAATTT	ATAGCGAGTC	CATCACCAAG	ACCTGCCGAA
1200	GTATTAGCTA	-AAATAATTAC	ATTAGCTCAT	AGAGCGACTC	TTAAAAAAAA	TGGTAACTAA
1260	TTTTTTTTT	TTTTGTATAA	ATAAAGCAGC	AATATAACTT	AAAATAAATT	ATTTTTTAA
1320	AGATGGGTTG	GAAAACGATG	TGTGCCGAGG	TTTTGGGAAG	TGTTTAGCAG	AAAAAAGTGT
1380	CCCGTACAGA	GCAGCATCGT	GCAAATATAG	GAAGAACACA	GGGAAGAAGT	GGGAAGGAGG

TCAGGCTGCA	ACCACGCCCC	GCGGAGATAG	TTAACGCGGC	CCACGTTGTG	CTATAGCCCG	1440
TCACTCTCGC	GGGCCTCTCC	AACCTCCAGT	TTTTTTTCTA	GCCCATCAGC	TGATACGGGG	1500
CCTTCCCCCC	ATGCAGGAGG	ATGGCCCGCC	ACGCGGTGTT	TTGGGCCGTT	CTCGCCGCGC	1560
GCGCCCGTGC	CGATCCGGGA	CTCATCCCAC	GTGCCGCCTC	GCCACCGCCG	CCGCCGCCGC	1620
TGCTGCTCCG	GCTGCCGGCT	GGACCTTCAC	GCTCACGCGC	TCTCCCCTGC	CCAACCACCA	1680
CGCAAACAAA	CACGAAGTTC	GCGCCGTCGA	CCGGCTCCCC	TCCTCCCCG	CGCGCATCGG	1740
ATCCCCCTAC	ATAAACCCTC	TCGCTCGCCA	TCGCCATGGC	AGCAACTCCC	CTCCTCCACT	1800
AGACCACCAT	GCACAGATCG	ATGGCCTCTC	AGGCGGTGGC	GCCCCTCCTC	CTCATCCTCA	1860
TGCTCGCGGC	GCCGCCGGG	GCCCCTCGG	CGGCGGTGCA	GTGCGGGCAG	GTGATGCAGC	1920
TGATGGCGCC	GTGCATGCCG	TACCTCGCCG	GCGCCCCCGG	GATGACGCCC	TACGGCATCT	1980
GCTGCGACAG	CCTCGGCGTG	CTCAACCGGA	TGGCCCCGGC	CCCCGCCGAC	CGCGTCGCCG	2040
TCTGCAACTG	CGTCAAGGAC	GCCGCCGCCG	GCTTCCCCGC	CGTCGACTTC	TCCCGCGCCT	2100
CCGCCCTCCC	CGCCGCCTGC	GGCCTCTCCA	TCAGCTTCAC	CATCGCCCCC	AACATGGACT	2160
GCAACCAGTA	AGTTCATTCA	TTCTTTCTTA	ACTCCAATTC	AATTTATCCA	TCACCTCGAC	2220
TTAAGCCTGA	TTAAACTTAA	CTTGTTCTTT	GCATGCTTGC	ACTATTGCAG	GGTTACAGAG	2280
GAACTGAGAA	TCTGAGAGCG	TGAGGAATCG	AGTTCATGTT	GCATTTATCA	TCAATCATCA	2340
TCGACTAGAT	CAATAAATCG	AGCAAAGCTT				2370

(2) INFORMATION FOR SEQ ID NO: 9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2407 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Oryza sativa
 - (B) STRAIN: Akihikari
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..2263
 - (D) OTHER INFORMATION: /label= PE1 /note= *sequence comprising anther specific promoter PE1*
- (ix) FEATURE:
 - (A) NAME/KEY: -

- (B) LOCATION: 2181..2187
- (D) OTHER INFORMATION: /label= TATA /note= "TATA Box"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2211
- (D) OTHER INFORMATION: /note= "transcription initiation site determined by primer extension"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2264..2266
- (D) OTHER INFORMATION: /label= ATG

/note= "ATG start of translation of El gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

60	ATATTGATTT	AGTTCTTTTC	TTCAAGTATC	GCTTTGTCAA	ATACTCACAT	TGATAGTGAC
120	TACATTTGGA	TTTTAGTAGG	CCATCAATTC	ATATTTCTTG	GAAAGTATAC	CTTAGTTGAT
180	ATCTGAGACT	TGATTTGCTT	GAGTCTCATC	CTCTTAACTG	CAGGGTTGAA	CACTAGTGGT
240	ACTTTGCTAC	AGTCCATGAC	GTGGACTGTC	ATGAGGCAAG	AAATCCTGTC	GGGTTTGTGC
300	ATGAATTATT	TCTCTTTCAA	GTATGTCCGT	CTTTTCCAGT	TTCTCGAAAT	TTCTATTAAG
360	ACAGGGCCTC	TTTTGTCTTC	TTCATTTAAC	CGGTGTACAT	GACAGCCTCG	TATATGTTCT
420	ATGTCGTCGC	GAACTTCTTG	CCTTCTTGTA	TTGGAATCAA	GTTGAGCAGA	TTGGTATTTT
480	ACTGGCAAAG	TTGGTACAAC	TCTTATATCT	TCAACTTCTG	AACTAGATGG	TACCCTTTGC
540	GAAAGTTTAG	TTGTCATTGT	AAATACAAAC	GTGAAGTAAG	CAAGAATCCT	TGTGCGCGCA
600	ATTGTTTTAT	CTTCTGTGTG	TCCTCAGATC	AAATTGTTTC	TCTTGACTCT	CTTTATATGA
660	AAGCTGCAAG	TAGACTTGTC	CAATATATAG	GAATACCTAC	TATTTATCTG	TAAAATTTAA
720	TAAACAATCT	CTTAATATCA	ATCCAACCAC	ACCAATAGAG	CGCCGACAAT	AACTTCCAAT
780	CATTATGAGG	AGCACATTAA	GAACAACAAT	ATTGAGTAGT	CCAGAACTAT	GATTGTTAGT
840	ATTTTAGCGC	AATCTGGTCA	TGATGCGTCT	TTCAATATTC	AACTCTGCAA	ATTATTGGCT
900	CATGTTTTTA	GGAACTGTTG	TGTTGGCACT	CCTTGGACAA	ATTGCACAAT	TCCAGAAAGA
960	TCTCTTCAGA	CCAGGAGAAA	TTATTATGTA	AAGGAGTAGA	TAACGTAGCA	CATCTCTTAT
1020	GTAATGGACG	ACGTTAGTTT	GATACAGTGT	GTAAAGAACA	ATGCAATGTC	TCCTTTCCAC
1080	CCTTGGAGGG	TTTCTGGGAT	GAGATGATGA	GGCATGTTCA	TTTCTCTGAA	GTCAATGCCA
1140	GTTATACTAC	AATGTCTTGC	TTTAGTACCT	TTAGTTGAGT	TCGGAAACAG	GCCCTGAAAT
1200	AAAGCTAATA	CACAGGAAAT	CTACCATCTC	GCTGAGTTTT	ATTTCTGTAA	GTGAAATGCC
1260	CAAGAATGGC	ACAAGCATGG	AATGAAGATC	CATTTGACCA	AGTGGTGCGG	CCTGTCCAAG
1320	ACCATATCAA	TCGAACAGGA	TTCTACTACA	TTATATTGTA	AGGAGCGGAA	AATCTGGCAA

TGTTGCCCCA	GCAAGGACCC	CCGCAGATAA	GTTCCTGTTC	TTCCACAGCA	GAATATCCGC	1380
AACTGCATAG	CTCCCAACAA	TGAAATCCAA	AACCACATCG	GCTCAGAGAG	AAGTTATGAT	1440
AAAAGGCACT	AATTCTGAAT	AATTTCCTAG	AAAGCGAATA	ATAATAGCAC	ACCTTGACCT	1500
CCACCAAGAA	GCTTGTGGAT	CGACTTGTGC	CCATGAAATG	GCATTCTGAC	ATTCTGGTCA	1560
CTGTCAGAAT	CTCTCGGAAA	ATGAGGAGGC	ATAGCTTCGT	GTGTGTATGT	GTGTGGGATA	1620
TTACGCTGCT	AAAACTTTGT	GTTTCTGATC	GATCTGGTTA	GAGAGCATCG	TCTTTATAAG	1680
CACTTAAAAA	TGGTAGTATA	ATCTCTCAAG	GAGCCTATAC	TGCCAAGGAA	AGGATAGCTT	1740
GGCCTGTGGG	GATTGAGCCG	TTGAAGGGAA	CAAACGAATA	CAGTTACCTT	ACCAGATGTT	1800
TGCCACGACA	TGGGCAACGT	CATTGCTAGA	CCAAGAAGGC	AAGAAGCAAA	GTTTAGCTGT	1860
CAAAAAAGAT	ATGCTAGAGG	CTTTCCAGAA	TATGTTCTAT	CTCAGCCAGA	CCAATGGGG	1920
CAAAATTTAC	TACTATTTGC	CATACATTAA	CCACGTAAAA	GTCCTACACT	CAACCTAACT	1980
GTTGAACGGT	CCTGTTCTGG	CCAACGGTGA	GAATGCACCT	AATGGACGGG	ACAACACTTC	2040
TTTCACCGTG	CTACTGCTAC	ATCCTGTAGA	CGGTGGACGC	GTGAGGTGCT	TTCGCCATGA	2100
CCGTCCTTGG	TTGTTGCAGT	CACTTGCGCA	CGCTTGCACC	GTGACTCACC	TGCCACATTG	2160
CCCCCGCCGT	CGCCGGCGCC	TACAAAAGCC	ACACACGCAC	GCCGGCCACG	ATAACCCATC	2220
CTAGCATCCC	GGTGTCCAGC	AAGAGATCCA	TCAAGCCGTC	GCGATGACGA	CGAGGCCTTC	2280
TGTTTTTTCC	ACCGTTGTCG	CGGCGATCGC	CATCGCCGCG	CTGCTGAGCA	GCCTCCTCCT	2340
CCTGCAGGCT	ACCCCGGCCG	CGGCCAGCGC	GAGGGCCTCG	AAGAAGGCTT	CGTGCGACCT	2400
GATGCAG						2407

(2) INFORMATION FOR SEQ ID NO: 10:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2784 base pairs (B) TYPE: nucleic acid

 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Zea mays
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..1179
 - (D) OTHER INFORMATION: /label= PCA55 /note= *region comprising the anther specific promoter and the leader sequence, PCA55*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 1072
 (D) OTHER INFORMATION: /label= TATA /note= "TATA Box"

(ix) FEATURE:

- (A) NAME/KEY: -(B) LOCATION: 1180..1596
- (D) OTHER INFORMATION: /note= "presumed coding sequence of corn CA55 gene*

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

TGGTATGCAT	CAATAGAGCC	GGAAGATGGT	CTGGAGTAAG	GACCTGGCAG	TGTGATACGG	60
GAACTTGACA	TCTGAATAGA	TATTCTCCCT	TGTCCCTCTG	GTAAAAAAA	CTGTTGTCAC	120
ATTTGCCTTC	GCTGTGACTT	GGATGTATCA	TGTATATCTT	TGACCATTGA	TATCTTGGTT	180
AATCAGACGG	TGCATTACAA	TCATGGCCTC	ATTCATATAG	GGTTTAGGGT	TACCACGATT	240
GGTTTGCATA	AGTAGTACCC	CTCCGTTTCA	AATTATGTCG	TATTTTGATT	TTTTAGATAC	300
ACTTTTTATA	TAATTTTTTA	TTTTAAATTA	GGTGTTTTAT	ATAATACGTA	TCTAAGTGTA	360
ТААТААААТА	TATGTATCTA	AAAGCTGTAA	TTTAGTATAA	ATTAGAATGG	TGTATATCTT	420
CAATGTATGA	CAAATAATTT	GAAATGGAGG	AGGGTATGAA	AAGCCAAAAC	CTCCTAGAAT	480
ATGGAATGGA	GGGAATACAT	ACAAATTCTT	TGCTTCAGTT	AAAAGAAACG	AGAAAAGGAG	540
GGGAATGGGG	AATCGTACTT	CAGTTTTTAC	GAGTTTTCAT	CAAACATGTA	TGCACGTCTT	600
CCCTTGGTTG	ATGCATCTTT	TTGGCAAATC	TTCGTTTAAT	TGCGGCTTCT	TTTTTATACC	660
GTTCGAAGGT	TTTCGTCGTC	AATGCTGAAA	CTCCACTTTC	ACCACCTTCG	GTTGCATCTG	720
CTTGCTTTCA	ATTCACCTCT	AATTAGTCCA	AGTGTTTCAT	TGGACGAAGG	TCCAAGTCCT	780
TCAGATCATC	TCAATTTTCT	TTGATCTGAA	ACAACAATTT	AAAACTGATT	TTGTTACCTT	840
GACCTGTCGA	AGACCTTCGA	ACGAACGGTA	CTGTAAAAAT	ACTGTACCTC	AGATTTGTGA	900
TTTCAATTCG	ATTCGGGTCT	CCTGGCTGGA	TGAAACCAAT	GCGAGAGAAG	AAGAAAAAAT	960
GTTGCATTAC	GCTCACTCGA	TCGGTTACGA	GCACGTAGTT	GGCGCCTGTC	ACCCAACCAA	1020
ACCAGTAGTT	GAGGCACGCC	CTGTTTGCTC	ACGATCACGA	ACGTACAGCA	CTATAAAACA	1080
CGCAGGGACT	GGAAAGCGAG	ATTTCACAGC	TCAAAGCAGC	CAAAACGCAG	AAGCTGCACT	1140
GCATATACAG	AAGATACATC	GAGCTAACTA	GCTGCAGCGA	TGTCTCGCTC	CTGCTGCGTC	1200
GCCGTGTCGG	TGCTTCTCGC	TGTCGCCGCG	ACAGCCAGCG	CCACCGCGCC	GGCATGGCTG	1260
CACGAGGAGC	AGCACCTCGA	GGAGGCCATG	GCCACGGGCC	CGCTGGTCGC	AGAGGGTGCG	1320
AGGGTGGCGC	CCTCCGCGTC	CACCTGGGCT	GCCGACAAGG	CGTCGCCGGC	GAGGCCGAGC	1380
GGCGGCATGG	CCACGCAGGG	CGACGACCAG	AGCTCGTCGG	GCGGCAGTGG	CAGCAGCGGT	1440

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	GAGCACGGCA	AGGCGGAGGG	CGAGAAGCAG	GGCAAGAGCT	GCCTCACCAA	GGAGGAGTGC	1500
	CACAAGAAGA	AGATGATCTG	TGGCAAGGGC	TGCACGCTCT	CGGCGCACAG	CAAGTGCGCC	1560
	GCCAAGTGCA	CCAAGTCCTG	TGTCCCCACC	TGCTAGGAGC	CGAGGCCGGA	GCTTGCCGGC	1620
	GGCGAGACCT	CGATCGATCG	AGTGCTTCAC	TTCACTTCTT	TGTTATAGTT	CTTGTGTGTT	1680
	GCCGTTGCGT	TGCGTTGCGT	AGACGAAGGG	AATAAGGAAG	GGTAATTGGA	TTACCTGTTC	1740
	CAGATCTCTG	TGTAAGCGTG	TTGTCGTGAC	AAGTCTTTTG	ATCCAGAGCG	AGGGATGGAT	1800
	AGATCGCGCT	CGCAGTTTTA	ATTGCAATGC	TAGTTCAATA	TGTGTGCATC	ATGTTGGCAA	1860
	CTACATAGTC	CAGATTCAAA	CCGAGATCGC	TGTTTAGCAT	GCCAGCACAA	TAATAACGGT	1920
	ACAATCATAT	TATATTTTAT	ACAAATGCAC	AATTTATCTC	TAGAGATGTC	AATGGGAAAT	1980
	TCCTCATCGG	GTTATATCAT	CTCAGACTCA	TCCCCATCAT	ATTTGATTCA	TCCTCATACT	2040
	CATCCTCATA	TCTATCATGA	GTGCAAAACT	CATTTCATAC	CCATCTCTAT	TTTGGTTTAG	2100
	GGTCTCCATC	ÇCTAATTAAG	GGATAACTAG	TACTAACAAC	TAGCACAAAC	TATCTAGATT	2160
	TCAGATATCA	CCACATTGAC	AAACAATCAT	CCATGAACTA	TGATCCATTC	ATCCATCCAT	2220
	СААААААТАА	ATCGGTATTT	CGAGAACGAT	AGAAGAAATG	AAGTCGGCTC	ACCTTTCTTG	2280
	GTCACCATTT	GAGTTTGTTG	GTGCCTGAGA	ATCCATGGTC	GTCATCGTCG	TCCTAGGGAT	2340
	CGGCGGTGCT	CCTCGTTGTT	GGTAAAGTCG	CCAGTGTGTA	GTGCTAGCGC	AACTGTCCAG	2400
	GCGTGCAACG	GTTGGCCGGC	TGGAAAGGGC	ATAGCGTATG	GCTGGTTATT	TTTAGGGTTT	2460
	TGTTTTTTA	CTAATCTGCT	AGTTGCCTTG	CCATGTTGTC	TTATTGGGCT	AGGATCTAGG	2520
	GCTTGTTACG	CTGCTGTGTT	GGGCTTGGTG	TCCGGTTCAG	CCTCAACTCA	TTCATACAAA	2580
	TCAGATTCAT	ACAAAACAGG	TATACACGTA	TGAAATATCC	ATGGATAATC	AGGTTCGAAT	2640
	TATTGTCCCC	TAAACCCATA	CACGTTTACC	CAATGGATGG	ATATTTTGTC	TCATATCCAT	2700
	ACACATGAGA	CGATTTTTGT	CCCATACCTG	TGCTCTAATA	GGAGAATTTC	TCTCGGGATA	2760
,	GCGAGTATCG	GATCCTCTAG	AGTC			•	2784

(2) INFORMATION FOR SEQ ID NO: 11:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 24 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:

WO 93/25695 PCT/EP93/01489

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(A) ORGANISM: oligonucleotide Zml3Oli2 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..24
(D) OTHER INFORMATION: /label= Zml3Oli2 /note= "oligonucleotide designated as Zml3Oli2" (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11: 24 GTGGATTGAA CGGGACTGAG TTGG (2) INFORMATION FOR SEO ID NO: 12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (iii) HYPOTHETICAL: NO (iii) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide Zml3Oli1 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..25 (D) OTHER INFORMATION: /label= Zm130li1 /note= "oligonucleotide designated as Zm130li1" (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12: AAGTCTCCAA GACTTTGGTT ATTCC 25 (2) INFORMATION FOR SEQ ID NO: 13: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (iii) HYPOTHETICAL: NO (iii) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANISM: Oligonucleotide Zml3Oli5 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..31 (D) OTHER INFORMATION: /label= Zml30li5

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/note= "oligonucleotide designated as Zm1301i5"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

GGATCCATGG TTGCCGCCGG GTGAATGTAC G

31

(2) INFORMATION FOR SEQ ID NO: 14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: oligonucleotide BXOL2
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..24
 - (D) OTHER INFORMATION: /label= BXOL2 /note= *oligonucleotide designated as BXOL2*
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

ACGGAAAACC TGAAGCACAC TCTC

24

(2) INFORMATION FOR SEQ ID NO: 15:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 49 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: oligonucleotide TA29SBXOL2
- (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..49
 - (D) OTHER INFORMATION: /label= TA29SBXOL2
 /note= "oligonucleotide designated as TA29SBXOL2"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

GTTTTTACTT AAAGAAATTA GCTACCATGA AAAAAGCAGT CATTAACGG

49

(2) INFORMATION FOR SEQ ID NO: 16:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(i

- i) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: oligonucleotide PTA290L5
 - (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: 1..27
 - (D) OTHER INFORMATION: /label= PTA290L5 /note= "oligonucleotide designated as PTA290L5"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

TGGCCATAAC TGAAATCAGG GTGAGAC

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- (2) INFORMATION FOR SEQ ID NO: 17:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 4808 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: EcoRI-HindIII fragment of plasmid pTS218
 - (ix) FEATURE:
 - (A) NAME/KEY: -
 - (B) LOCATION: complement (18..401)
 - (D) OTHER INFORMATION: /label= 3'nos
 /note= "3' regulatory sequence containing the
 polyadenylation site derived from Agrobacterium
 T-DNA nopaline synthase gene"
 - (ix) FEATURE:
 - (A) NAME/KEY: --
 - (B) LOCATION: complement (402..737)
 - (D) OTHER INFORMATION: /label= barnase /note= "coding region of the barnase gene of Bacillus amyloliquefaciens"
 - (ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (738..1944)
- (D) OTHER INFORMATION: /label= PZM13
 /note= "promoter region of the Zm13 gene of Zea mays"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (1945..2281)
- (D) OTHER INFORMATION: /label= 3'nos

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (2282..2554)
- (D) OTHER INFORMATION: /label= barstar

/note= "coding region of the barstar gene of Bacillus amyloliquefaciens"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (2555..3099)
- (D) OTHER INFORMATION: /label= PTA29

/note= *promoter region of the TA29 gene of Nicotiana tabacum*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 3100..3932
- (D) OTHER INFORMATION: /label= 35S3

/note= ""35S3" promoter sequence derived from cauliflower mosaic virus isolate CabbB-JI"

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 3933..4484
- (D) OTHER INFORMATION: /label= bar
 /note= *coding region of the phosphinothricin
 acetyltransferase gene*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 4485..4763
- (D) OTHER INFORMATION: /label= 3'nos

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 2333..2356
- (D) OTHER INFORMATION: /label= BXOL2
 /note= *region corresponding to oligonucleotide
 BXOL2*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (2538..2586)
 (D) OTHER INFORMATION: /label= TA29SBXOL2
- /note= *region complementary to oligonucleotide TA29SBXOL2*

(ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: complement (2800..2823)
- (D) OTHER INFORMATION: /label= PTA290L5

/note= "region complementary to part of oligonucleotide PTA29OL5"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

GAATTCGAGC	TCGGTACCCG	GGGATCTTCC	CGATCTAGTA	ACATAGATGA	CACCGCGCGC	60
GATAATTTAT	CCTAGTTTGC	GCGCTATATT	TTGTTTTCTA	TCGCGTATTA	AATGTATAAT	120
TGCGGGACTC	TAATCATAAA	AACCCATCTC	ATAAATAACG	TCATGCATTA	CATGTTAATT	180
ATTACATGCT	TAACGTAATT	CAACAGAAAT	TATATGATAA	TCATCGCAAG	ACCGGCAACA	240
GGATTCAATC	TTAAGAAACT	TTATTGCCAA	ATGTTTGAAC	GATCTGCTTC	GGATCCTCTA	300
GAGNNNNCCG	GAAAGTGAAA	TTGACCGATC	AGAGTTTGAA	GAAAAATTTA	TTACACACTT	360
TATGTAAAGC	TGAAAAAAAC	GGCCTCCGCA	GGAAGCCGTT	TTTTTCGTTA	TCTGATTTTT	420
GTAAAGGTCT	GATAATGGTC	CGTTGTTTTG	TAAATCAGCC	AGTCGCTTGA	GTAAAGAATC	480
CGGTCTGAAT	TTCTGAAGCC	TGATGTATAG	TTAATATCCG	CTTCACGCCA	TGTTCGTCCG	540
CTTTTGCCCG	GGAGTTTGCC	TTCCCTGTTT	GAGAAGATGT	CTCCGCCGAT	GCTTTTCCCC	. 600
GGAGCGACGT	CTGCAAGGTT	CCCTTTTGAT	GCCACCCAGC	CGAGGGCTTG	TGCTTCTGAT	660
TTTGTAATGT	AATTATCAGG	TAGCTTATGA	TATGTCTGAA	GATAATCCGC	AACCCCGTCA	720
AACGTGTTGA	TAACCGGTAC	CATGGTTGCC	GCCGGGTGAA	TGTACGTGTT	TTCCCCTCCC	780
CCCTTGTGGA	TGTCGGAGGA	AAAGGGCGGG	ACCTTTCCTT	ATTATTTGTG	TGAGGGAGGG	840
AGGGTGAGAC	GAGGGTGGCA	AATCTGGCCT	GGTTTCGTAA	CGCAGCCTGT	GGTGTCCTTA	900
AATAATCTGC	GAGACCCGAA	GATTTGTCTG	GCTGCCCGTA	GAAATAGTGG	CCGACGGCCA	960
GAAAGCCGTT	TTGACCTAGT	TTTGTGGCAT	GGGGAGGTGA	AAAATATGTC	GAATTTGGTT	1020
GAGTTTTTCA	GCGGTCCACA	AAAGTTGGAT	TTGCGACTTC	TAATCTTCTA	TGCTGCTATG	1080
TAAAACTAAA	TAAGACGCGC	TCAACAAAAC	GGTATCAGTT	AAATGTATCC	ATAAGTGAAT	1140
AGTATATAGA	ATCACTAAAT	GGCAGCTTCT	CATATTTAGC	ATAGTACTTG	GACAAATGTC	1200
ACGCAAGATG	TATTATAAAA	ACACATCGGT	GTTGAGTGTT	GATACAATAA	AATCTCAATA	1260
GTTGTTTTAA	TCTCAATCCA	TGTGGATTGA	ACGGGACTGA	GTTGGTTTAA	ATCCCTAGTA	1320
AGTCAAAATT	ATTTCTAATT	TTATCTAAAT	CTCATCTAAT	CCACGTGGAA	TAGGAATAAC	1380
CAAACTGTTG	GAGACTTGTT	CTCAAGTGCT	ATGAGTTAAG	AACAAGGCAA	CATAGAAAAT	1440
ATTAATCGTT	AAGGTCCTTC	GTCCTTCGAA	GCATTATTTC	CCTTAGGATA	TAATGGTTTA	1500
CGGACGAAGG	TTATGAAGGA	CGTACCTTCA	TAAATTCATT	AAACAATGAC	GAAGGATGAA	1560
ATATAAAGAA	TATAAAAGAC	AACATGAACA	ATTATATATT	ATTATTAGGT	ACAAACAGAA	1620
ATATCGTTGA	ATTACAAGTG	TACCTTCAAT	AGGAATGAGA	TGACAGTACA	AGCGTGACGC	1680
AAAAAGCGAA	TGCCAAGTCA	GCGTGAACAG	TACGGGAATA	CTGTTCACCT	ATTTATAGGC	1740.

ACGGGACGTA	GCCTGTGCAA	AATTACATTA	ATGCCCTTTA	CACTTAATA	TAAACCTATA	1800
GTAATCTGTT	GAGGTCTAAA	TAGCCTTTTC	ATCTTTAAGT	CGGTTTCAAC	TGCTGCTGTC	1860
TTGCCGAAGC	TTTCCTGCTI	ACACCTTAGG	CGCTTCACCA	ACCTTCGTAT	TATTCTGGTC	1920
TACTGTGATG	CCTGACTTGA	GTCCGAAGAT	GGGGATCTTC	CCGATCTAGT	AACATAGATG	1980
ACACCGCGCG	CGATAATTTA	TCCTAGTTTG	CGCGCTATAT	TTTGTTTTCT	ATCGCGTATT	2040
AAATGTATAA	TTGCGGGACT	CTAATCATAA	AAACCCATCT	CATAAATAAC	GTCATGCATT	2100
ACATGTTAAT	TATTACATGC	TTAACGTAAT	TCAACAGAAA	TTATATGATA	ATCATCGCAA	2160
GACCGGCAAC	AGGATTCAAT	CTTAAGAAAC	TTTATTGCCA	AATGTTTGAA	CGATCTGCTT	2220
CGGATCCTCT	AGACCAAGCT	AGCTTGCGGG	TTTGTGTTTC	CATATTGTTC	ATCTCCCATT	2280
GATCGTATTA	AGAAAGTATG	ATGGTGATGT	CGCAGCCTTC	CGCTTTCGCT	TCACGGAAAA	2340
CCTGAAGCAC	ACTCTCGGCG	CCATTTTCAG	TCAGCTGCTT	GCTTTGTTCA	AACTGCCTCC	2400
ATTCCAAAAC	GAGCGGGTAC	TCCACCCATC	CGGTCAGACA	ATCCCATAAA	GCGTCCAGGT	2460
TTTCACCGTA	GTATTCCGGA	AGGGCAAGCT	CCTTTTTCAA	TGTCTGGTGG	AGGTCGCTGA	2520
TACTTCTGAT	TTGTTCCCCG	TTAATGACTG	CTTTTTTCAT	GGTAGCTAAT	TTCTTTAAGT	2580
AAAAACTTTG	ATTTGAGTGA	TGATGTTGTA	CTGTTACACT	TGCACCACAA	GGGCATATAT	2640
AGAGCACAAG	ACATACACAA	CAACTTGCAA	AACTAACTTT	TGTTGGAGCA	TTTCGAGGAA	2700
AATGGGGAGT	AGCAGGCTAA	TCTGAGGGTA	ACATTAAGGT	TTCATGTATT	AATTTGTTGC	2760
AAACATGGAC	TTAGTGTGAG	GAAAAAGTAC	CAAAATTTTG	TCTCACCCTG	ATTTCAGTTA	2820
TGGAAATTAC	ATTATGAAGC	TGTGCTAGAG	AAGATGTTTA	TTCTAGTCCA	GCCACCCACC	2880
TTATGCAAGT	CTGCTTTTAG	CTTGATTCAA	AAACTGATTT	AATTTACATT	GCTAAATGTG	2940
CATACTTCGA	GCCTATGTCG	CTTTAATTCG	AGTAGGATGT	ATATATTAGT	ACATAAAAAA	3000
TCATGTTTGA	ATCATCTTTC	ATAAAGTGAC	AAGTCAATTG	TCCCTTCTTG	TTTGGCACTA	3060
TATTCAATCT	GTTAATGCAA	ATTATCCAGT	TATACTTAGC	TAGATCCTAC	GCAGCAGGTC	3120
TCATCAAGAC	GATCTACCCG	AGTAACAATC	TCCAGGAGAT	CAAATACCTT	CCCAAGAAGG	3180 -
TTAAAGATGC	AGTCAAAAGA	TTCAGGACTA	ATTGCATCAA	GAACACAGAG	AAAGACATAT	3240
TTCTCAAGAT	CAGAAGTACT	ATTCCAGTAT	GGACGATTCA	AGGCTTGCTT	CATAAACCAA	3300
GGCAAGTAAT	AGAGATTGGA	GTCTCTAAAA	AGGTAGTTCC	TACTGAATCT	AAGGCCATGC	3360
ATGGAGTCTA	AGATTCAAAT	CGAGGATCTA	ACAGAACTCG	CCGTGAAGAC	TGGCGAACAG	3420
TTCATACAGA	GTCTTTTACG	ACTCAATGAC	AAGAAGAAAA	TCTTCGTCAA	CATGGTGGAG	3480
CACGACACTC	TGGTCTACTC	CAAAAATGTC	AAAGATACAG	TCTCAGAAGA	CCAAAGGGCT	3540
ATTGAGACTT	TTCAACAAAG	GATAATTTCG	GGAAACCTCC	TCGGATTCCA	TTGCCCAGCT	3600

				,		
ATCTGTCACT	TCATCGAAAG	GACAGTAGAA	AAGGAAGGTG	GCTCCTACAA	ATGCCATCAT	3660
TGCGATAAAG	GAAAGGCTAT	CATTCAAGAT	GCCTCTGCCG	ACAGTGGTCC	CAAAGATGGA	3720
CCCCCACCCA	CGAGGAGCAT	CGTGGAAAAA	GAAGACGTTC	CAACCACGTC	TTCAAAGCAA	3780
GTGGATTGAT	GTGACATCTC	CACTGACGTA	AGGGATGACG	CACAATCCCA	CTATCCTTCG	3840
CAAGACCCTT	CCTCTATATA	AGGAAGTTCA	TTTCATTTGG	AGAGGACACG	CTGAAATCAC	3900
CAGTCTCTCT	CTATAAATCT	ATCTCTCTCT	CTATAACCAT	GGACCCAGAA	CGACGCCCGG	3960
CCGACATCCG	CCGTGCCACC	GAGGCGGACA	TGCCGGCGGT	CTGCACCATC	GTCAACCACT	4020
ACATCGAGAC	AAGCACGGTC	AACTTCCGTA	CCGAGCCGCA	GGAACCGCAG	GAGTGGACGG	4080
ACGACCTCGT	CCGTCTGCGG	GAGCGCTATC	CCTGGCTCGT	CGCCGAGGTG	GACGGCGAGG	4140
TCGCCGGCAT	CGCCTACGCG	GGCCCCTGGA	AGGCACGCAA	CGCCTACGAC	TGGACGGCCG	4200
AGTCGACCGT	GTACGTCTCC	CCCCGCCACC	AGCGGACGGG	ACTGGGCTCC	ACGCTCTACA	4260
CCCACCTGCT	GAAGTCCCTG	GAGGCACAGG	GCTTCAAGAG	CGTGGTCGCT	GTCATCGGGC	4320
TGCCCAACGA	CCCGAGCGTG	CGCATGCACG	AGGCGCTCGG	ATATGCCCCC	CGCGGCATGC	4380
TGCGGGCGGC	CGGCTTCAAG	CACGGGAACT	GGCATGACGT	GGGTTTCTGG	CAGCTGGACT	4440
TCAGCCTGCC	GGTACCGCCC	CGTCCGGTCC	TGCCCGTCAC	CGAGATCTGA	TCTCACGCGT	4500
CTAGGATCCG	AAGCAGATCG	TTCAAACATT	TGGCAATAAA	GTTTCTTAAG	ATTGAATCCT	4560
GTTGCCGGTC	TTGCGATGAT	TATCATATAA	TTTCTGTTGA	ATTACGTTAA	GCATGTAATA	4620
ATTAACATGT	AATGCATGAC	GTTATTTATG	AGATGGGTTT	TTATGATTAG	AGTCCCGCAA	4680
TTATACATTT	AATACGCGAT	AGAAAACAAA	ATATAGCGCG	CAAACTAGGA	TAAATTATCG	4740
CGCGCGGTGT	CATCTATGTT	ACTAGATCGG	GAAGATCCTC	TAGAGTCGAC	CTGCAGGCAT	4800
GCAAGCTT						4808

FIGURE 1

1. <u>Action</u>: Transform corn embryos (e.g. H99) with malesterility gene S, linked to herbicide resistance gene <u>bar</u> (Example 5)

Result: transformed plants with genotype S/s

2. <u>Action</u>: Transform corn embryos (e.g. H99) with fertility-restorer gene R (Example 6)

Result: transformed plants with genotype R/r

3. Action : Transform corn embryos (e.g; H99) with maintainer gene P (Example 3)

Result : transformed plants with genotype P/p

4. Action : Cross S/s,r/r x s/s,R/r. Select offspring for presence of both S and R genes by means of PCR. Result : plants with genotype S/s,R/r

5. Action : Self selected plants of 4 (optional)
Result : Progeny plants with 9 different genotypes

gamete ♀ ♂→ ↓	S,R	s,r	s,R	s,r
S,R	S/S,R/R	S/S,R/r	S/s,R/R	S/s,R/r
S,r	S/S,R/r	S/S,r/r**	S/s,R/r	S/s,r/r**
s,R	S/s,R/R	S/s,R/r	s/s,R/R	s/s,R/r
s,r	S/s,R/r	S/s,r/r**	s/s,R/r	s/s,r/r

[&]quot; male-sterile plants

6. <u>Action</u>: self male-fertile progeny plants of 5 (Optional) Result:

6.1. Self of S/S,R/R : 100 % male-fertile plants

6.2 Self of S/s,R/r : Same progeny as in 5

13/16 male-fertile plants with 4/13 herbicide sensitive

FIGURE 1 (continued 1)

6.3 Self of S/S,R/r : Progeny as follows :

gamete ♀ ♂→ ↓	S,R	s,r
S,R	S/S,R/R	S/S,R/r
S,r	S/S,R/r	S/S,r/r**

[&]quot; male-sterile plants

Thus: 3/4 male-fertile plants, 0% herbicide sensitive All male-sterile plants are of genotype S/S,r/r

7. Action : Cross

?: P/p (from 3) $x \neq 3$: S/s,R/r (from 4)

this equals in fact

 $9: s/s, r/r, P/p \times \sigma: S/s, R/r, p/p$

Result: Progeny with the following genotypes

gamete ♀ ♂→ ↓	S,R,p	S,r,p	s,R,p	s,r,p
s,r,P	S/s,R/r,P/p	S/s,r/r,P/p	s/s,R/r,P/p	s/s,r/r,P/p
s,r,p	S/s,R/r,p/p	S/s,r/r,p/p	s/s,R/r,p/p	s/s,r/r,p/p

8. <u>Action</u>: From offspring of 7, select plants with genotype S/s,r/r,P/p by screening, by means of PCR and/or Southern blotting, for presence of S and P gene and absence of R gene.

Result : plants with genotype S/s,P/p

FIGURE 1 (continued 2)

9. Action: Self plants with genotype S/s,P/p (from 8)
Result: progeny with the following genotypes

gamete ♀ ♂→	 S,P	S,p	s,P	s,p
S, P	S/S,P/P	S/S,P/p	S/s,P/P	S/s,P/p
S,p	S/S,P/p	S/S,p/p"	S/s,P/p	S/s,p/p**
s,P	S/s,P/P	S/s,P/p	s/s,P/P	s/s,P/p
s,p	S/s,P/p	S/s,p/p**	8/3.P/p	s/s,p/p

" male-sterile plants
Shaded genotypes cannot develop because male
gametes (pollen) are killed off by expression of
the maintainer gene P.

10. Action : self male fertile plants of 9. Result

10.1. Self of s/s,P/p : 100 % male-fertile plants Self of s/s,p/p : 100 % male-fertile plants

10.2 Self of S/s,P/p : Same progeny as in 9
5/8 male-fertile plants with
2/5 herbicide sensitive

10.3 Self of S/S,P/p : Progeny as follows :

gamete ♀ ♂→ ↓	S, P	S,p	
S,P	5/5,P/P	S/S,P/p	
S,p	S/S, P/p	S/S,p/p"	

" male-sterile plants
Shaded genotypes cannot develop because male
gametes (pollen) are killed off by
expression of the maintainer gene P.

FINAL RESULT

- 1/2 male-fertile plants, 0% herbicide sensitive. All these plants are maintainer plants
- 1/2 male sterile plants. All homozygous for the malesterility gene S.

CLAIMS

- 1. A cell of a maintainer plant, the nuclear genome of which contains: 1) at a first locus, a malesterility genotype in homozygous condition; and 2) at a second locus, a maintainer gene in heterozygous condition; said first and second loci preferably being unlinked; said maintainer gene being a foreign DNA sequence, preferably a foreign chimaeric DNA sequence, including:
- a) a fertility-restorer gene that comprises:
- a fertility-restorer DNA encoding a restorer RNA and/or protein or polypeptide which, when produced or overproduced in cells, preferably stamen cells, of said plant, prevents phenotypic expression of said nuclear male-sterility genotype which would render said plant male-sterile in the absence of said restorer RNA, protein or polypeptide in said stamen cells and
- ii) a restorer promoter capable of directing expression of said fertility-restorer DNA at least in said cells, preferably said stamen cells, so that said phenotypic expression of said nuclear male-sterility genotype is prevented, said fertility-restorer DNA being in the same transcriptional unit as, and under the control of, said restorer promoter and
- b) a pollen-lethality gene that is selectively expressed in microspores and/or pollen of said plant to prevent the production of functional pollen and that comprises:
- iii) a pollen-lethality DNA coding for a pollenlethality RNA and/or protein or polypetide that, when produced or overproduced in said microspores and/or pollen, significantly disrupts the metabolism, functioning and/or

- development of said microspores and/or pollen and
- iv) a pollen-specific promoter capable of directing expression of said pollen-lethality DNA selectively in said microspores and/or pollen said pollen-lethality DNA being in the same transcriptional unit as, and under the control of, said pollen-specific promoter.
- 2. The cell of claim 1, in which said maintainer gene also contains, preferably in said second locus, a first marker gene which comprises:
- v) a first marker DNA encoding a first marker RNA and/or protein or polypeptide which, when present at least in a first specific tissue or specific cells of said plant, renders said plant easily separable from other plants which do not contain said first marker RNA, protein or polypeptide at least in said first specific tissue or specific cells and
- vi) a first marker promoter capable of directing expression of said first marker DNA at least in said first specific tissue or specific cells, said first marker DNA being in the same transcriptional unit as, and under the control of, said first marker promoter.
- 3. The cell of claim 1 or 2, in which said malesterility genotype is foreign to said plant and is a male-sterility gene that is a foreign DNA sequence, preferably a foreign chimaeric DNA sequence, comprising:
- 1) a male-sterility DNA encoding a sterility RNA and/or protein or polypeptide which, when produced or overproduced in said stamen cells of said plant in the absence of said restorer RNA, protein or polypeptide, significantly disturbs the metabolism,

functioning and/or development of said stamen cells and

- 2) a sterility promoter capable of directing expression of said male-sterility DNA selectively in said stamen cells, said male-sterility DNA being in the same transcriptional unit as, and under the control of, said sterility promoter.
- 4. The cell of claim 3 in which said male-sterility genotype also contains, preferably in said first locus, a second marker gene which comprises:
- 3) a second marker DNA encoding a second marker RNA and/or protein or polypeptide which, when present at least in a second specific tissue or specific cells of said plant, renders said plant easily separable from other plants which do not contain said second marker RNA, protein or polypeptide at least in said second specific tissue or specific cells and
- 4) a second marker promoter capable of directing expression of said second marker DNA at least in said specific tissue or specific cells, said second marker DNA being in the same transcriptional unit as, and under the control of, said second marker promoter.
- 5. The cell of claim 3 or 4, in which said malesterility DNA encodes Barnase.
- 6. The cell of any one of claims 3 to 5, in which said sterility promoter is selected among the TA29 promoter of <u>Nicotiana tabacum</u> and the promoters of SEQ ID nos. 7 to 10.
- 7. The cell of claim 5 or 6, in which said restorer DNA encodes Barstar.
- 8. The cell of any one of claims 1 to 7, in which said restorer promoter is selected among the TA29 promoter of <u>Nicotiana tabacum</u> and the promoters of SEQ ID nos. 7 to 10.

- 9. The cell of claim 1 or 2, in which said malesterility genotype is endogenous to said plant and is homozygous for a recessive allele.
- 10. The cell of claim 9, in which said fertilityrestorer gene is the dominant allele of the
 endogenous male-sterility genotype, preferably under
 the control of its natural promoter.
- 11. The cell of any one of claims 1 to 10, in which said pollen-lethality DNA encodes a ribonuclease, preferably RNAse T1 or Barnase.
- 12. The cell of any one of claims 1 to 11, in which said pollen-specific promoter is the Zm13 promoter of SEQ ID no. 1.
- The cell of any one of claims 2 to 12, in which said first marker DNA or said second marker DNA is: an herbicide resistance gene, particularly an sfr or sfrv gene; a gene encoding a modified target enzyme for an herbicide having lower affinity for the herbicide, particularly modified а enolpyruvylshikimate-3 phosphate synthase as a target for glyphosate or a modified glutamine synthetase as a target for a glutamine synthetase inhibitor such as phosphinotricine; a gene encoding a protein which confers resistance to the herbicide sulfonylurea; a gene encoding a protein which confers resistance to the herbicide, bromoxynil; a gene encoding a protein which confers resistance to the herbicide 2,4 D; a gene encoding a protein or a polypeptide conferring a color to at least said first or second specific tissue or specific cells, particularly the Al gene or the glucuronidase gene; a gene encoding a protein or a polypeptide conferring a stress tolerance to said plant, particularly a gene encoding Mn-superoxide dismutase; a gene encoding a protein or a polypeptide conferring a disease or pest resistance, particularly

- a gene encoding a <u>Bacillus thuringiensis</u> endotoxin that confers insect resistance; or a gene encoding a bactericidal peptide that confers a bacterial resistance.
- 14. The cell of any one of claims 2 to 13, in which said first marker promoter or said second marker promoter is: a constitutive promoter, particularly a 35S promoter, a 35S'3 promoter, a PNOS promoter or a POCS promoter; a wound-inducible promoter, particularly a TR1' or TR2' promoter; a promoter which directs gene expression selectively in plant tissue having photosynthetic activity, particularly an SSU promoter; or a promoter which directs gene expression selectively in leaf cells, petal cells or seed cells, particularly seed coat cells.
- 15. The cell of any one of claims 2 to 12, wherein said first marker DNA and said second marker DNA are different.
- 16. A cell of a plant, the nuclear genome of which contains the maintainer gene of any one of claims 1 to 14.
- 17. A plant cell culture consisting essentially of the plant cells of any one of claims 1 to 16.
- 18. A plant, particularly corn, oilseed rape, wheat, rice, sunflower, sugarbeet, tomato, lettuce, peppers, sorghum, soybean, pea, alfalfa, grasses, clovers, carrot, cabbages, leek, onion, tobacco, petunia, cacao and citrus, more particularly corn, oilseed rape, wheat and rice, consisting essentially of the plant cells of any one of claims 1 to 16.
- 19. A seed of the plant of claim 18.
- 20. The maintainer gene of any one of claims 1 to 14.

- 21. A vector for transforming a cell of a plant, comprising the maintainer gene of any one of claims 1 to 14, particularly pTS210.
- 22. A method to maintain a homogeneous population of male-sterile plants or their seed, the nuclear genome of which contain, at said first locus of any one of claims 1, 3-6 and 9, said male-sterility genotype of any one of claims 1, 3-6 and 9 in homozygous condition; said method comprising the step of crossing said male-sterile plants with the plant of claim 18.

International Application No

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